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(54) Title: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR ANALYSIS OF GENE EXPRESSION IN HUMAN BREAST AND HBL 100 CELLS

(57) Abstract: A single exon nucleic acid microarray comprising a plurality of single exon nucleic acid probes for measuring gene expression in a sample derived from human HBL 100 cells is described. Also described are single exon nucleic acid probes expressed in the HBL 100 cells and their use in methods for detecting gene expression.

HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR ANALYSIS OF GENE EXPRESSION IN HUMAN BREAST AND HBL 100 CELLS

#### 5 CROSS REFERENCE TO RELATED APPLICATIONS

The present application is a continuation-in-part of U.S. patent application serial nos. 09/632,366, filed August 3, 2000 and 09/608,408, filed June 30, 2000; claims the

10 benefit under 35 U.S.C. s 119(e) of U.S.provisional patent application serial nos. 60/236,359, filed September 27, 2000, 60/234,687, filed September 21, 2000, 60/207,456, filed May 26, 2000, and 60/180,312, filed February 4, 2000; and further claims the benefit under 35 U.S.C. s 119(a) of

15 UK patent application no. 0024263.6, filed October 4, 2000, the disclosures of which are incorporated herein by reference in their entireties.

REFERENCE TO SEQUENCE LISTING AND INCORPORATION BY
20 REFERENCE THEREOF

The present application includes a Sequence Listing in electronic format, filed pursuant to PCT Administrative Instructions 801 - 806 on a single CD-R disc, in triplicate, containing a file named pto\_HBL100.txt, created 24 January 2001, having 11,029,597 bytes. The Sequence Listing contained in said file on said disc is incorporated herein by reference in its entirety.

#### 30 Field of the Invention

The present invention relates to genome-derived single exon microarrays useful for verifying the expression of regions of genomic DNA predicted to encode protein. In particular, the present invention relates to unique genome-

derived single exon nucleic acid probes expressed in human HBL 100 cells and single exon nucleic acid microarrays that include such probes.

# 5 Background of the Invention

For almost two decades following the invention of general techniques for nucleic acid sequencing, Sanger et al., Proc. Natl. Acad. Sci. USA 70(4):1209-13 (1973); Gilbert et al., Proc. Natl. Acad. Sci. USA 70(12):3581-4 (1973), these techniques were used principally as tools to further the understanding of proteins — known or suspected — about which a basic foundation of biological knowledge had already been built. In many cases, the cloning effort that preceded sequence identification had been both informed and directed by that antecedent biological understanding.

For example, the cloning of the T cell receptor for antigen was predicated upon its known or suspected cell type-specific expression, by its suspected membrane

20 association, and by the predicted assembly of its gene via T cell-specific somatic recombination. Subsequent sequencing efforts at once confirmed and extended

understanding of this family of proteins. Hedrick et al.,

Nature 308(5955):153-8 (1984).

More recently, however, the development of high throughput sequencing methods and devices, in concert with large public and private undertakings to sequence the human and other genomes, has altered this investigational paradigm: today, sequence information often precedes understanding of the basic biology of the encoded protein product.

One of the approaches to large-scale sequencing is predicated upon the proposition that expressed sequences — that is, those accessible through isolation of mRNA — are of greatest initial interest. This "expressed

sequence tag" ("EST") approach has already yielded vast amounts of sequence data (see for example Adams et al., Science 252:1651 (1991); Williamson, Drug Discov. Today 4:115 (1999)). For nucleic acids sequenced by this approach, often the only biological information that is known a priori with any certainty is the likelihood of biologic expression itself. By virtue of the species and tissue from which the mRNA had originally been obtained, most such sequences are also annotated with the identity of the species and at least one tissue in which expression appears likely.

More recently, the pace of genomic sequencing has accelerated dramatically. When genomic DNA serves as the initial substrate for sequencing efforts, expression cannot be presumed; often the only a priori biological information about the sequence includes the species and chromosome (and perhaps chromosomal map location) of origin.

accumulation by directed, EST, and genomic sequencing

approaches — and in particular, with the accumulation of sequence information from multiple genera, from multiple species within genera, and from multiple individuals within a species — there is an increasing need for methods that rapidly and effectively permit the functions of nucleic sequences to be elucidated. And as such functional information accumulates, there is a further need for methods of storing such functional information in meaningful and useful relationship to the sequence itself; that is, there is an increasing need for means and apparatus for annotating raw sequence data with known or predicted functional information.

Although the increase in the pace of genomic sequencing is due in large part to technological changes in sequencing strategies and instrumentation, Service, Science 280:995 (1998); Pennisi, Science 283: 1822-1823 (1999),

there is an important functional motivation as well.

While it was understood that the EST approach would rarely be able to yield sequence information about the noncoding portions of the genome, it now also appears the EST approach is capable of capturing only a fraction of a genome's actual expression complexity.

For example, when the C. elegans genome was fully sequenced, gene prediction algorithms identified over 19,000 potential genes, of which only 7,000 had been found 10 by EST sequencing. C. elegans Sequencing Consortium, Science 282:2012 (1998). Analogously, the recently completed sequence of chromosome 2 of Arabidopsis predicts over 4000 genes, Lin et al., Nature, 402:761 (1999), of which only about 6% had previously been identified via EST 15 sequencing efforts. Although the human genome has the greatest depth of EST coverage, it is still woefully short of surrendering all of its genes. One recent estimate suggests that the human genome contains more than 146,000 genes, which would at this point leave greater than half of 20 the genes undiscovered. It is now predicted that many genes, perhaps 20 to 50%, will only be found by genomic sequencing.

There is, therefore, a need for methods that permit the functional regions of genomic sequence — and 25 most importantly, but not exclusively, regions that function to encode genes — to be identified.

Much of the coding sequence of the human genome is not homologous to known genes, making detection of open reading frames ("ORFs") and predictions of gene function difficult. Computational methods exist for predicting coding regions in eukaryotic genomes. Gene prediction programs such as GRAIL and GRAIL II, Uberbacher et al., Proc. Natl. Acad. Sci. USA 88(24):11261-5 (1991); Xu et al., Genet. Eng. 16:241-53 (1994); Uberbacher et al.,

Methods Enzymol. 266:259-81 (1996); GENEFINDER, Solovyev et

al., Nucl. Acids. Res. 22:5156-63 (1994); Solovyev et al., Ismb 5:294-302 (1997); and GENESCAN, Burge et al., J. Mol. Biol. 268:78-94 (1997), predict many putative genes without known homology or function. Such programs are known,

5 however, to give high false positive rates. Burset et al., Genomics 34:353-367 (1996). Using a consensus obtained by a plurality of such programs is known to increase the reliability of calling exons from genomic sequence. Ansari-Lari et al., Genome Res. 8(1):29-40 (1998)

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Identification of functional genes from genomic data remains, however, an imperfect art. For example, in reporting the full sequence of human chromosome 21, the Chromosome 21 Mapping and Sequencing Consortium reports that prior bioinformatic estimates of human gene number may 15 need to be revised substantially downwards. 405:311-199 (2000); Reeves, Nature 405:283-284 (2000).

Thus, there is a need for methods and apparatus that permit the functions of the regions identified bioinformatically - and specifically, that permit the 20 expression of regions predicted to encode protein - readily to be confirmed experimentally.

Recently, the development of nucleic acid microarrays has made possible the automated and highly parallel measurement of gene expression. Reviewed in 25 Schena (ed.), DNA Microarrays : A Practical Approach (Practical Approach Series), Oxford University Press (1999) (ISBN: 0199637768); Nature Genet. 21(1)(suppl):1 - 60 (1999); Schena (ed.), Microarray Biochip: Tools and Technology, Eaton Publishing Company/BioTechniques Books 30 Division (2000) (ISBN: 1881299376).

It is common for microarrays to be derived from cDNA/EST libraries, either from those previously described in the literature, such as those from the I.M.A.G.E. consortium, Lennon et al., Genomics 33(1):151-2 (1996), or 35 from the construction of "problem specific" libraries

targeted at a particular biological question, R.S. Thomas et al., Cancer Res. (in press). Such microarrays by definition can measure expression only of those genes found in EST libraries, and thus have not been useful as probes for genes discovered solely by genomic sequencing.

The utility of using whole genome nucleic acid microarrays to answer certain biological questions has been demonstrated for the yeast Saccharomyces cerevisiae. De Risi et al., Science 278:680 (1997). The vast majority of yeast nuclear genes, approximately 95% however, are single exon genes, i.e., lack introns, Lopez et al., RNA 5:1135-1137 (1999); Goffeau et al., Science 274:563-67 (1996), permitting coding regions more readily to be identified. Whole genome nucleic acid microarrays have not generally been used to probe gene expression from more complex eukaryotic genomes, and in particular from those averaging more than one intron per gene.

Diseases of the breast are a significant cause of human morbidity and mortality. Increasingly, genetic

20 factors are being found that contribute to predisposition, onset, and/or aggressiveness of most, if not all, of these diseases. Although mutations in single genes have been identified as causative for some diseases of the breast, for the most part these disorders are believed to have polygenic etiologies. There is a need for methods and apparatus that permit prediction, diagnosis and prognosis of diseases of the human breast, particularly those diseases with polygenic etiology.

#### 30 Summary of the Invention

The present invention solves these and other problems in the art by providing methods and apparatus for predicting, confirming, and displaying functional information derived from genomic sequence. The present

invention also provides apparatus for verifying the expression of putative genes identified within genomic sequence.

In particular, the invention provides novel

5 genome-derived single exon nucleic acid microarrays useful
for verifying the expression of putative genes identified
within genomic sequence.

The present invention also provides compositions and kits for the ready production of nucleic acids

10 identical in sequence to, or substantially identical in sequence to, probes on the genome-derived single exon microarrays of the present invention.

Accordingly, in a first aspect of the invention, there is provided a spatially-addressable set of single

15 exon nucleic acid probes for measuring gene expression in a sample derived from human Breast, comprising a plurality of single exon nucleic acid probes according to any one of the nucleotide sequences set out in SEQ ID NOs: 1 - 5,074 or a complementary sequence, or a portion of such a sequence.

20 By plurality is meant at least two, suitably at least 20, most suitably at least 100, preferably at least 1000 and, most preferably, upto 5000.

In one embodiment of the first aspect, each of said plurality of probes is separately and addressably amplifiable.

In an alternative embodiment, each of said plurality of probes is separately and addressably isolatable from said plurality.

In a preferred embodiment, each of said plurality
of probes is amplifiable using at least one common primer.
Preferably, each of said plurality of probes is amplifiable
using a first and a second common primer.

In yet another embodiment, said set of single exon nucleic acid probes comprises between 50 - 20,000 probes, for example, 50 - 5000.

Suitably, said set of single exon nucleic acid probes comprises at least 50 - 1000 discrete single exon nucleic acid probes having a sequence as set out in any of SEQ ID NOS.: 1 - 10,058 or a complimentary sequence, or a portion of such a sequence.

Preferably, the average length of the single exon nucleic acid probes is between 200 and 500 bp. It is preferred that the average length should be at least 200bp, suitably at least 250bp, most suitably at least 300bp, preferably at least 400bp and, most preferably, 500 bp.

In another embodiment, the single exon nucleic acid probes lack prokaryotic and bacteriophage vector sequence. It is preferred that at least 50%, suitably at least 60%, most suitably at least 70%, preferably at least 75%, more preferably at least 80, 85, 90, 95 or 99% of said single exon nucleic acid probes lack prokaryotic and bacteriophage vector sequence.

In another preferred embodiment, said single exon nucleic acid lack homopolymeric stretches of A or T. It is preferred that at least 50%, suitably at least 60%, most suitably at least 70%, preferably at least 75%, more preferably at least 80, 85, 90, 95 or 99% of said single exon nucleic acid probes lack homopolymeric stretches of A or T.

25 Preferably, a spatially-addressable set of single exon nucleic acid probes in accordance with the first aspect of the invention is is addressably disposed upon a substrate.

Suitable substrates include a filter membrane

30 which may, preferably, be nitrocellulose or nylon. The
nylon may preferably, be positively-charged. Other suitable
substrates include glass, amorphous silicon, crystalline
silicon, and plastic. Further suitable materials include
polymethylacrylic, polyethylene, polypropylene,

35 polyacrylate, polymethylmethacrylate, polyvinylchloride,

polytetrafluoroethylene, polystyrene, polycarbonate, polyacetal, polysulfone, celluloseacetate, cellulosenitrate, nitrocellulose, and mixtures thereof.

In a second aspect of the invention, there is 5 provided a microarray comprising a spatially addressable set of single exon nucleic acid probes in accordance with the first aspect of the invention.

In one embodiment, a genome-derived single-exon microarray is packaged together with such an ordered set of 10 amplifiable probes corresponding to the probes, or one or more subsets of probes, thereon. In alternative embodiments, the ordered set of amplifiable probes is packaged separately from the genome-derived single exon microarray.

In another aspect, the invention provides genomederived single exon nucleic acid probes useful for gene expression analysis, and particularly for gene expression analysis by microarray. In particular embodiments of this aspect, the present invention provides human single-exon 20 probes that include specifically-hybridizable fragments of SEQ ID Nos. 5,075 - 10,058, wherein the fragment hybridizes at high stringency to an expressed human gene. particular embodiments, the invention provides single exon probes comprising SEQ ID Nos. 1 - 5,074.

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25 Accordingly, in a third aspect of the invention, there is provided a single exon nucleic acid probe for measuring human gene expression in a sample derived from human Breast which is a nucleic acid molecule comprising a nucleotide sequence as set out in any of SEQ ID NOs.: 1 -30 5,074 or a complementary sequence or a fragment thereof wherein said probe hybridizes at high stringency to a nucleic acid expressed in the human Breast.

In one embodiment, a single exon nucleic acid probe in accordance with the third aspect comprises a 35 nucleotide sequence as set out in any of SEQ ID NOs.: 5,075

- 10,058 or a complementary sequence or a fragment thereof.

In a fourth aspect of the invention, there is provided a single exon nucleic acid probe for measuring human gene expression in a sample derived from human Breast 5 which is a nucleic acid molecule having a sequence encoding a peptide comprising a peptide sequence as set out in any of SEQ ID NOs.: 10,059 - 15,009 or a complementary sequence or a fragment thereof wherein said probe hybridizes at high stringency to a nucleic acid expressed in the human Breast.

Preferably, a single exon nucleic acid probe in accordance with the third or fourth aspects of the invention comprises between at least 15 and 50 contiguous nucleotides of said SEQ ID NO:. It is preferred that the single exon nucleic acid probe comprises at least 15, 15 suitably at least 20, more suitably at least 25 or preferably at least 50 contiguous nucleotides of said SEQ ID NO:.

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In another preferred embodiment, a single exon nucleic acid probe in accordance with the third or fourth 20 aspects of the invention is between 3kb and 25kb in length. It is preferred that said probe is no more than 3kb, suitably no more than 5kb, more suitably no more than 10kb, preferably 15kb, more preferably 20kb or, most preferably, no more than 20kb in length.

25 Preferably, a single exon nucleic acid probe in accordance with either the fifth or sixth aspect of the invention is DNA, preferably single-stranded DNA, RNA or PNA.

In another embodiment of either the third or 30 fourth aspect of the invention, a single exon nucleic acid probe is detectably labeled. Suitable detectable labels include a radionuclide, a fluorescent label or a first member of a specific binding pair. Suitable fluorescent labels include dyes such as cyanine dyes, preferably Cy3 and Cy5 although other suitable dyes will be known to those

skilled in the art.

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In a particularly preferred embodiment, a single exon nucleic acid probe in accordance with either the third or fourth aspect of the invention lacks prokaryotic and bacteriophage vector sequence. In yet another embodiment, a single exon nucleic acid probe in accordance with either the third or fourth aspect of the invention lacks homopolymeric stretches of A or T.

In a fifth aspect of the invention, there is provided an amplifiable nucleic acid composition, comprising:

the single exon nucleic acid probe in accordance with either of the third or fourth aspects of the invention; and at least one nucleic acid primer;

wherein said at least one primer is sufficient to prime enzymatic amplification of said probe.

In an sixth aspect of the invention, there is provided a method of measuring gene expression in a sample derived from human Breast, comprising:

contacting the single exon microarray in accordance with the second aspect of the invention, with a first collection of detectably labeled nucleic acids, said first-collection of nucleic acids derived from mRNA of human Breast; and then

25 measuring the label detectably bound to each probe of said microarray.

In a seventh aspect of the invention, there is provided a method of identifying exons in a eukaryotic genome, comprising:

algorithmically predicting at least one exon from genomic sequence of said eukaryote; and then

detecting specific hybridization of detectably labeled nucleic acids to a single exon probe,

wherein said detectably labeled nucleic acids are 35 derived from mRNA from the Breast of said eukaryote, said

probe is a single exon probe having a fragment identical in sequence to, or complementary in sequence to, said predicted exon, said probe is included within a single exon microarray in accordance with the first aspect of the invention, and said fragment is selectively hybridizable at high stringency.

In a eighth aspect of the invention, there is provided a method of assigning exons to a single gene, comprising:

identifying a plurality of exons from genomic sequence in accordance with the seventh aspect of the invention; and then

measuring the expression of each of said exons in a plurality of tissues and/or cell types using

15 hybridization to single exon microarrays having a probe with said exon,

wherein a common pattern of expression of said exons in said plurality of tissues and/or cell types indicates that the exons should be assigned to a single gene.

In an ninth aspect of the invention, there is provided a nucleic acid sequence as set out in any of SEQ

ID NOs: 1 -- 10,058 wherein said sequence encodes a peptide.

In a tenth aspect of the invention, there is 25 provided a peptide encoded by a sequence comprising a sequence as set out in any of SEQ ID NOs: 5,075 - 10,058, or a complementary sequence or coding portion thereof.

In a preferred embodiment, a peptide may be encoded by a sequence comprising a sequence set out in any of SEQ ID NOS.: 1 -5,074.

In a further aspect, the invention provides peptides comprising an amino acid sequence translated from the DNA fragments, said amino acid sequences comprising SEQ ID NOS.: 10,059 - 15,009.

Accordingly in a eleventh aspect of the invention

there is provided a peptide comprising a sequence as set out in any of SEQ ID NOs: 10,059 - 15,009, or fragment thereof.

In another aspect, the invention provides means

for displaying annotated sequence, and in particular, for
displaying sequence annotated according to the methods and
apparatus of the present invention. Further, such display
can be used as a preferred graphical user interface for
electronic search, query, and analysis of such annotated
sequence.

#### Detailed Description of the Invention

#### 15 Definitions

As used herein, the term "microarray" and phrase "nucleic acid microarray" refer to a substrate-bound collection of plural nucleic acids, hybridization to each of the plurality of bound nucleic acids being separately detectable. The substrate can be solid or porous, planar or non-planar, unitary or distributed.

"nucleic acid microarray" include all the devices so called in Schena (ed.), DNA Microarrays: A Practical Approach

(Practical Approach Series), Oxford University Press (1999)

(ISBN: 0199637768); Nature Genet. 21(1) (suppl):1 - 60

(1999); and Schena (ed.), Microarray Biochip: Tools and Technology, Eaton Publishing Company/BioTechniques Books Division (2000) (ISBN: 1881299376). As so defined, the

term "microarray" and phrase "nucleic acid microarray" further include substrate-bound collections of plural nucleic acids in which the nucleic acids are distributably disposed on a plurality of beads, rather than on a unitary planar substrate, as is described, inter alia, in Brenner et al., Proc. Natl. Acad. Sci. USA 97(4):166501670 (2000);

in such case, the term "microarray" and phrase "nucleic acid microarray" refer to the plurality of beads in aggregate.

As used herein with respect to a nucleic acid 5 microarray, the term "probe" refers to the nucleic acid that is, or is intended to be, bound to the substrate; in such context, the term "target" thus refers to nucleic acid intended to be bound thereto by Watson-Crick complementarity. As used herein with respect to solution 10 phase hybridization, the term "probe" refers to the nucleic acid of known sequence that is detectably labeled.

As used herein, the expression "probe comprising SEQ ID NO.", and variants thereof, intends a nucleic acid probe, at least a portion of which probe has either (i) the 15 sequence directly as given in the referenced SEQ ID NO., or (ii) a sequence complementary to the sequence as given in the referenced SEQ ID NO., the choice as between sequence directly as given and complement thereof dictated by the requirement that the probe hybridize to mRNA.

As used herein, the term "open reading frame" and the equivalent acronym "ORF" refer to that portion of an exon that can be translated in its entirety into a sequence of contiguous amino acids i.e. a nucleic acid sequence that, in at least one reading frame, does not possess stop 25 codons; the term does not require that the ORF encode the entirety of a natural protein.

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As used herein, the term "amplicon" refers to a PCR product amplified from human genomic DNA, containing the predicted exon.

As used herein the term "exon" refers to the consensus prediction of the various exon and gene predicting algorithms i.e. a nucleic acid sequence bioinformatically predicted to encode a portion of a natural protein.

As used herein, the term "peptide" refers to a

sequence of amino acids. The sequences referred to as PEPTIDE SEQ ID NOS.: are the predicted peptide sequences that would be translated from one of the exons, or a portion thereof set out in exon SEQ ID NOS.:. The codons encoding the peptide are wholly contained within the exon.

As used herein, a "portions" of a defined nucleotide sequence or sequences can be and, preferably, are fragments unique to that sequence or to one or a combination of those sequences. A fragment unique to a nucleic acid molecule is one that is a signature for the larger nucleic acid molecule.

As used herein, the phrase "expression of a probe" and its linguistic variants means that the ORF present within the probe, or its complement, is present within a target mRNA.

As used herein, "stringent conditions" refers to parameters well known to those skilled in the art. When a nucleic acid molecule is said to be hybridisable to another of a given sequence under "stringent conditions" it is meant that it is homologous to the given sequence.

As used herein, the phrase "specific binding pair" intends a pair of molecules that bind to one another with high specificity. Binding pairs are said to exhibit specific binding when they exhibit avidity of at least 10<sup>7</sup>, preferably at least 10<sup>8</sup>, more preferably at least 10<sup>9</sup> liters/mole. Nonlimiting examples of specific binding pairs are: antibody and antigen; biotin and avidin; and biotin and streptavidin.

As used herein with respect to the visual display of annotated genomic sequence, the term "rectangle" means any geometric shape that has at least a first and a second border, wherein the first and second borders each are capable of mapping uniquely to a point of another visual object of the display.

As used herein, a "Mondrian" means a visual

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display in which a single genomic sequence is annotated with predicted and experimentally confirmed functional information.

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## Brief Description of the Drawings

The present invention is further illustrated with reference to the following non-limiting figures and examples in which:

FIG. 1 illustrates a process for predicting functional regions from genomic sequence, confirming the functional activity of such regions experimentally, and associating and displaying the data so obtained in meaningful and useful relationship to the original sequence data:

FIG. 2 further elaborates that portion of the process schematized in FIG. 1 for predicting functional regions from genomic sequence;

FIG. 3 illustrates a Mondrian visual display;

FIG. 4 presents a Mondrian showing a hypothetical annotated genomic sequence;

FIG. 5 is a histogram showing the distribution of ORF length and PCR products as obtained, with ORF length shown in black and PCR product length shown in dotted lines;

FIG. 6 is a histogram showing the distribution, among exons predicted according to the methods described, of expression as measured using simultaneous two color hybridization to a genome-derived single exon microarray. The graph shows the number of sequence-verified products that were either not expressed ("0"), expressed in one or more but not all tested tissues ("1" - "9"), or expressed in all tissues tested ("10");

FIG. 7 is a pictorial representation of the

expression of verified sequences that showed expression with signal intensity greater than 3 in at least one tissue, with: FIG. 7A showing the expression as measured by microarray hybridization in each of the 10 measured tissues, and the expression as measured "bioinformatically" by query of EST, NR and SwissProt databases; with FIG. 7B showing the legend for display of physical expression (ratio) in FIG. 7A; and with FIG. 7C showing the legend for scoring EST hits as depicted in FIG. 7A;

signal intensity for arrayed sequences that were identical to sequences in existing EST, NR and SwissProt databases or that were dissimilar (unknown), where black denotes the signal intensity for all sequence-verified products with a BLAST Expect ("E") value of greater than 1e-30 (1 x 10<sup>-30</sup>) ("unknown") and a dotted line denotes sequence-verified spots with a BLAST expect ("E") value of less than 1e-30 (1 x 10<sup>-30</sup>) ("known");

FIG. 9 presents a Mondrian of BAC AC008172 (bases 20 25,000 to 130,000), containing the carbamyl phosphate synthetase gene (AF154830.1); and

FIG. 10 is a Mondrian of BAC A049839.

Methods and Apparatus for Predicting, Confirming,
Annotating, and Displaying Functional Regions From Genomic
Sequence Data

FIG. 1 is a flow chart illustrating in broad
outline a process for predicting functional regions from
genomic sequence, confirming and characterizing the
functional activity of such regions experimentally, and
then associating and displaying the information so obtained
in meaningful and useful relationship to the original
sequence data.

The initial input into process 10 of the present invention is drawn from one or more databases 100 containing genomic sequence data. Because genomic sequence is usually obtained from subgenomic fragments, the sequence data typically will be stored in a series of records corresponding to these subgenomic sequenced fragments. Some fragments will have been catenated to form larger contiguous sequences ("contigs"); others will not. A finite percentage of sequence data in the database will typically be erroneous, consisting inter alia of vector sequence, sequence created from aberrant cloning events, sequence of artificial polylinkers, and sequence that was erroneously read.

Each sequence record in database 100 will

minimally contain as annotation a unique sequence
identifier (accession number), and will typically be
annotated further to identify the date of accession,
species of origin, and depositor. Because database 100 can
contain nongenomic sequence, each sequence will typically

be annotated further to permit query for genomic sequence.
Chromosomal origin, optionally with map location, can also
be present. Data can be, and over time increasingly will
be, further annotated with additional information, in part
through use of the present invention, as described below.

Annotation can be present within the data records, in

25 Annotation can be present within the data records, in information external to database 100 and linked to the records thereto, or through a combination of the two.

Databases useful as genomic sequence database 100 in the present invention include GenBank, and particularly include several divisions thereof, including the htgs(draft), NT (nucleotide, command line), and NR (nonredundant) divisions. GenBank is produced by the National Institutes of Health and is maintained by the National Center for Biotechnology Information (NCBI).

35 Databases of genomic sequence from species other than

human, such as mouse, rat, Arabidopsis, C. elegans, C. brigsii, Drosophila, zebra fish, and other higher eukaryotic organisms will also prove useful as genomic sequence database 100.

Genomic sequence obtained by query of genomic 5 sequence database 100 is then input into one or more processes 200 for identification of regions therein that are predicted to have a biological function as specified by the user. Such functions include, but are not limited to, 10 encoding protein, regulating transcription, regulating message transport after transcription into mRNA, regulating message splicing after transcription into mRNA, of regulating message degradation after transcription into mRNA, and the like. Other functions include directing 15 somatic recombination events, contributing to chromosomal stability or movement, contributing to allelic exclusion or X chromosome inactivation, and the like.

The particular genomic sequence to be input into process 200 will depend upon the function for which 20 relevant sequence is to be identified as well as upon the approach chosen for such identification. Process step 200 can be iterated to identify different functions within a given genomic region. In such case, the input often will be different for the several iterations.

Sequences predicted to have the requisite function by process 200 are then input into process 300, where a subset of the input sequences suitable for experimental confirmation is identified. Experimental confirmation can involve physical and/or bioinformatic 30 assay. Where the subsequent experimental assay is bioinformatic, rather than physical, there are fewer constraints on the sequences that can be tested, and in this latter case therefore process 300 can output the entirety of the input sequence.

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The subset of sequences output from process 300 35

is then used in process 400 for experimental verification and characterization of the function predicted in process 200, which experimental verification can, and often will, include both physical and bioinformatic assay.

process 500 annotates the sequence data with the functional information obtained in the physical and/or bioinformatic assays of process 400. Such annotation can be done using any technique that usefully relates the functional information to the sequence, as, for example, by incorporating the functional data into the sequence data record itself, by linking records in a hierarchical or relational database, by linking to external databases, by a combination thereof, or by other means well known within the database arts. The data can even be submitted for incorporation into databases maintained by others, such as GenBank, which is maintained by NCBI.

As further noted in FIG. 1, additional annotation can be input into process 500 from external sources 600.

The annotated data is then displayed in process 800, either before, concomitantly with, or after optional storage 700 on nontransient media, such as magnetic disk, optical disc, magnetooptical disk, flash memory, or the like.

FIG. 1 shows that the experimental data output

25 from process 400 can be used in each preceding step of
process 10: e.g., facilitating identification of functional
sequences in process 200, facilitating identification of an
experimentally suitable subset thereof in process 300, and
facilitating creation of physical and/or informational
30 substrates for, and performance of subsequent assay, of
functional sequences in process 400.

Information from each step can be passed directly to the succeeding process, or stored in permanent or interim form prior to passage to the succeeding process.

35 Often, data will be stored after each, or at least a

plurality, of such process steps. Any or all process steps can be automated.

FIG. 2 further elaborates the prediction of functional sequence within genomic sequence according to process 200.

Genomic sequence database 100 is first queried 20 for genomic sequence.

The sequence required to be returned by query 20 will depend, in the first instance, upon the function to be identified.

For example, genomic sequences that function to encode protein can be identified inter alia using gene prediction approaches, comparative sequence analysis approaches, or combinations of the two. In gene prediction analysis, sequence from one genome is input into process 200 where at least one, preferably a plurality, of algorithmic methods are applied to identify putative coding regions. In comparative sequence analysis, by contrast, corresponding, e.g., syntenic, sequence from a plurality of sources, typically a plurality of species, is input into process 200, where at least one, possibly a plurality, of algorithmic methods are applied to compare the sequences and identify regions of least variability.

The exact content of query 20 will also depend
upon the database queried. For example, if the database
contains both genomic and nongenomic sequence, perhaps
derived from multiple species, and the function to be
determined is protein coding regions in human genomic
sequence, the query will accordingly require that the
sequence returned be genomic and derived from humans.

Query 20 can also incorporate criteria that compel return of sequence that meets operative requirements of the subsequent analytical method. Alternatively, or in addition, such operative criteria can be enforced in subsequent preprocess step 24.

For example, if the function sought to be identified is protein coding, query 20 can incorporate criteria that return from genomic sequence database 100 only those sequences present within contigs sufficiently long as to have obviated substantial fragmentation of any given exon among a plurality of separate sequence fragments.

Such criteria can, for example, consist of a required minimal individual genomic sequence fragment

length, such as 10 kb, more typically 20 kb, 30 kb, 40kb, and preferably 50 kb or more, as well as an optional further or alternative requirement that sequence from any given clone, such as a bacterial artificial chromosome ("BAC"), be presented in no more than a finite maximal number of fragments, such as no more than 20 separate pieces, more typically no more than about 10 - 12 fragments.

Results using the present invention have shown that genomic sequence from bacterial artificial chromosomes 20 (BACs) is sufficient for gene prediction analysis according to the present invention if the sequence is at least 50 kb in length, and if additionally the sequence from any given BAC is presented in fewer than 15, and preferably fewer than 10, fragments. Accordingly, query 20 can incorporate 25 a requirement that data accessioned from BAC sequencing be in fewer than 15, preferably fewer than 10, fragments.

An additional criterion that can be incorporated into the query can be the date, or range of dates, of sequence accession. Although the process has been described above as if genomic sequence database 100 were static, it is of course understood that the genomic sequence databases need not be static, and indeed are typically updated on a frequent, even hourly, basis. Thus, as further described in Examples 1 and 2, infra, it is possible to query the database for newly added sequence,

either newly added after an absolute date, or newly added relative to a prior analysis performed using the methods and apparatus of the present invention. In this way, the process herein described can incorporate a dynamic, 5 temporal component.

One utility of such temporal limitation is to identify, from newly accessioned genomic sequence, the presence of novel genes, particularly those not previously identified by EST sequencing (or other sequencing efforts 10 that are similarly based upon gene expression). As further described in Example 1, such an approach has shown that newly accessioned human genomic sequence, when analyzed for sequences that function to encode protein, readily identifies genes that are novel over those in existing EST 15 and other expression databases. This makes the methods of the present invention extremely powerful gene discovery tools. And as would be appreciated, such gene discovery can be performed using genomic sequence from species other than human.

If query 20 incorporates multiple criteria, such. as above-described, the multiple criteria can be performed as a series of separate queries or as a single query, depending in part upon the query language, the complexity of the query, and other considerations well known in the 25 database arts.

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If query 20 returns no genomic sequence meeting the query criteria, the negative result can be reported by process 22, and process 200 (and indeed, entire process 10) ended 23, as shown. Alternatively, or in addition to 30 report and termination of the initial inquiry, a new query 20 can be generated that takes into account the initial negative result.

When query 20 returns sequence meeting the query criteria, the returned sequence is then passed to optional 35 preprocessing 24, suitable and specific for the desired

analytical approach and the particular analytical methods thereof to be used in process 25.

Preprocessing 24 can include processes suitable for many approaches and methods thereof, as well as processes specifically suited for the intended subsequent analysis.

Preprocessing 24 suitable for most approaches and methods will include elimination of sequence irrelevant to, or that would interfere with, the subsequent analysis.

10 Such sequence includes repetitive sequence, such as Alu repeats and LINE elements, vector sequence, artificial sequence, such as artificial polylinkers, and the like.

Such removal can readily be performed by identification and subsequent masking of the undesired sequence.

Identification can be effected by comparing the genomic sequence returned by query 20 with public or private databases containing known repetitive sequence, vector sequence, artificial sequence, and other artifactual sequence. Such comparison can readily be done using programs well known in the art, such as CROSS\_MATCH, or by proprietary sequence comparison programs the engineering of which is well within the skill in the art.

Alternatively, or in addition, undesirable, including artifactual, sequence can be identified

25 algorithmically without comparison to external databases and thereafter removed. For example, synthetic polylinker sequence can be identified by an algorithm that identifies a significantly higher than average density of known restriction sites. As another example, vector sequence can be identified by algorithms that identify nucleotide or codon usage at variance with that of the bulk of the genomic sequence.

Once identified, undesired sequence can be removed. Removal can usefully be done by masking the undesired sequence as, for example, by converting the

specific nucleotide references to one that is unrecognized by the subsequent bioinformatic algorithms, such as "X". Alternatively, but at present less preferred, the undesired sequence can be excised from the returned genomic sequence, leaving gaps.

Preprocessing 24 can further include selection from among duplicative sequences of that one sequence of highest quality. Higher quality can be measured as a lower percentage of, fewest number of, or least densely clustered 10 occurrence of ambiguous nucleotides, defined as those nucleotides that are identified in the genomic sequence using symbols indicating ambiguity. Higher quality can also or alternatively be valued by presence in the longest contig.

Preprocessing 24 can, and often will, also include formatting of the data as specifically appropriate for passage to the analytical algorithms of process 25. Such formatting can and typically will include, inter alia, addition of a unique sequence identifier, either derived 20 from the original accession number in genomic sequence database 100, or newly applied, and can further include additional annotation. Formatting can include conversion from one to another sequence listing standard, such as conversion to or from FASTA or the like, depending upon the 25 input expected by the subsequent process.

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Preprocessing, which can be optional depending upon the function desired to be identified and the informational requirements of the methods for effecting such identification, is followed by sequence processing 25 30 where sequences with the desired function are identified within the genomic sequence.

As mentioned above, such functions can include, but are not limited to, encoding protein, regulating transcription, regulating message transport after 35 transcription into mRNA, regulating message splicing after

transcription, of regulating message degradation, and the like. Other functions include directing somatic recombination events, contributing to chromosomal stability or movement, contributing to allelic exclusion or X chromosome inactivation, or the like.

The methods of the present invention are particularly useful for gene discovery, that is, for identifying, from genomic sequence, regions that function to encode genes, and in a particularly useful embodiment, for identifying regions that function to encode genes not hitherto identified by expression-based or directed cloning and sequencing. In conjunction with verification using the novel single exon microarrays of the present invention, as further described below, the methods herein described become powerful gene discovery tools.

Accordingly, in a preferred embodiment of the present invention, process 25 is used to identify putative coding regions. Two preferred approaches in process 25 for identifying sequence that encodes putative genes are gene prediction and comparative sequence analysis.

Gene prediction can be performed using any of a number of algorithmic methods, embodied in one or more software programs, that identify open reading frames (ORFs) using a variety of heuristics, such as GRAIL, DICTION, and GENEFINDER. Comparative sequence analysis similarly can be performed using any of a variety of known programs that identify regions with lower sequence variability.

As further described in Example 1, below, gene finding software programs yield a range of results. For the newly accessioned human genomic sequence input in Example 1, for example, GRAIL identified the greatest percentage of genomic sequence as putative coding region, 2% of the data analyzed; GENEFINDER was second, calling 1%; and DICTION yielded the least putative coding region, with 0.8% of genomic sequence called as coding region.

Increased reliability can be obtained when consensus is required among several such methods. Although discussed herein particularly with respect to exon calling, consensus among methods will in general increase 5 reliability of predicting other functions as well.

Thus, as indicated by query 26, sequence processing 25, optionally with preprocessing 24, can be repeated with a different method, with consensus among such iterations determined and reported in process 27.

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Process 27 compares the several outputs for a given input genomic sequence and identifies consensus among the separately reported results. The consensus itself, as well as the sequence meeting that consensus, is then stored in process 29a, displayed in process 29b, and/or output to 15 process 300 for subsequent identification of a subset thereof suitable for assay.

Multiple levels of consensus can be calculated and reported by process 27. For example, as further described in Example 1, infra, process 27 can report 20 consensus as between all specific pairs of methods of gene prediction, as consensus among any one or more of the pairs of methods of gene prediction, or as among all of the gene prediction algorithms used. Thus, in Example 1, process 27 reported that GRAIL and GENEFINDER programs agreed on 0.7% 25 of genomic sequence, that GRAIL and DICTION agreed on 0.5% of genomic sequence, and that the three programs together agreed on 0.25% of the data analyzed. Put another way, 0.25% of the genomic sequence was identified by all three of the programs as containing putative coding region.

Furthermore, consensus can be required among different approaches to identifying a chosen function.

For example, if the function desired to be identified is coding of protein sequence, and a first used approach to exon calling is gene prediction, the process 35 can be repeated on the same input sequence, or subset

thereof, with another approach, such as comparative sequence analysis. In such a case, where comparative sequence analysis follows gene prediction, the comparison can be performed not only on genomic nucleic acid sequence, but additionally or alternatively can be performed on the predicted amino acid sequence translated from the ORFs prior identified by the gene prediction approach.

Although shown as an iterative process, the multiple analyses required to achieve consensus can be done in series, in parallel, or some combination thereof.

Predicted functional sequence, optionally representing a consensus among a plurality of methods and approaches for determination thereof, is passed to process 300 for identification of a subset thereof for functional assay.

In the preferred embodiment of the methods of the present invention, wherein the function sought to be identified is protein coding, process 300 is used to identify a subset thereof suitable for experimental verification by physical and/or bioinformatic approaches.

For example, putative ORFs identified in process 200 can be classified, or binned, bioinformatically into putative genes. This binning can be based inter alia upon consideration of the average number of exons/gene in the species chosen for analysis, upon density of exons that have been called on the genomic sequence, and other empirical rules. Thereafter, one or more among the genespecific ORFs can be chosen for subsequent use in gene expression assay.

Where such subsequent gene expression assay uses amplified nucleic acid, considerations such as desired amplicon length, primer synthesis requirements, putative exon length, sequence GC content, existence of possible secondary structure, and the like can be used to identify and select those ORFs that appear most likely successfully

to amplify. Where subsequent gene expression assay relies upon nucleic acid hybridization, whether or not using amplified product, further considerations involving hybridization stringency can be applied to identify that 5 subset of sequences that will most readily permit sequencespecific discrimination at a chosen hybridization and wash stringency. One particular such consideration is avoidance of putative exons that span repetitive sequence; such sequence can hybridize spuriously to nonspecific message, reducing specific signal in the hybridization.

For bioinformatic assay, there are fewer constraints on the sequences that can be tested experimentally, and in this latter case therefore process 300 can output the entirety of the input sequence.

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The subset of sequences identified by process 300 as suitable for use in assay is then used in process 400 to create the physical and/or informational substrate for experimental verification of the predictions made in process 200, and thereafter to assay those substrates.

As mentioned, the methods of the present invention are particularly useful for identifying potential coding regions within genomic sequence. In a preferred embodiment of process 400, therefore, the expression of the sequences predicted to encode protein is verified. 25 combination of the predictive and experimental methods provides a powerful gene discovery engine.

Thus, in another aspect, the present invention provides methods and apparatus for verifying the expression of putative genes identified within genomic sequence. 30 particular, the invention provides a novel method of verifying gene expression in which expression of predicted ORFs is measured and confirmed using a novel type of nucleic acid microarray, the genome-derived single exon nucleic acid microarrays of the present invention.

Putative ORFs as predicted by a consensus of gene

calling, particularly gene prediction, algorithms in process 200, and as further identified as suitable by process 300, are amplified from genomic DNA using the polymerase chain reaction (PCR). Although PCR is conveniently used, other amplification approaches can also be used.

Amplification schemes can be designed to capture the entirety of each predicted ORF in an amplicon with minimal additional (that is, intronic or intergenic)

10 sequence. Because ORFs predicted from human genomic sequence using the methods of the present invention differ in length, such an approach results in amplicons of varying length.

However, most predicted ORFs are shorter than 500 bp in length, and although amplicons of at least about 100 or 200 base pairs can be immobilized as probes on nucleic acid microarrays, early experimental results using the methods of the present invention have suggested that longer amplicons, at least about 400 or 500 base pairs, are more effective. Furthermore, certain advantages derive from application to the microarray of amplicons of defined size.

Therefore, amplification schemes can alternatively, and preferably, be designed to amplify regions of defined size, preferably at least about 300, 400 or 500 bp, centered about each predicted ORF. Such an approach results in a population of amplicons of limited size diversity, but that typically contain intronic and/or intergenic nucleic acid in addition to putative ORF.

Conversely, somewhat fewer than 10% of ORFs

predicted from human genomic sequence according to the
methods of the present invention exceed 500 bp in length.

Portions of such extended ORFs, preferably at least about
300,400 or 500 bp in length, can be amplified. However, it
has been discovered that the percentage success at
amplifying pieces of such ORFs is low, and that such

putative exons are more effectively amplified when larger fragments, at least about 1000 or 1500 bp, and even as large as 2000 bp are amplified.

The putative ORFs selected in process 300 are

thus input into one or more primer design programs, such as
PRIMER3 (available online for use at
http://www-genome.wi.mit.edu/cgi-bin/primer/), with a goal
of amplifying at least about 500 base pairs of genomic
sequence centered within or about ORFs predicted to be no
more than about 500 bp, or at least about 1000 - 1500 bp of
genomic sequence for ORFs predicted to exceed 500 bp in
length, and the primers synthesized by standard techniques.
Primers with the requisite sequences can be purchased
commercially or synthesized by standard techniques.

15 Conveniently, a first predetermined sequence can be added commonly to the ORF-specific 5' primer and a second, typically different, predetermined sequence commonly added to each 3' ORF-unique primer. This serves to immortalize the amplicon, that is, serves to permit 20 further amplification of any amplicon using a single set of primers complementary respectively to the common 5' and common 3' sequence elements. The presence of these

"universal" priming sequences further facilitates later sequence verification, providing a sequence common to all 25 amplicons at which to prime sequencing reactions. The common 5' and 3' sequences further serve to add a cloning site should any of the ORFs warrant further study.

Such predetermined sequence is usefully at least about 10, 12 or 15 nt in length, and usually does not exceed about 25 nt in length. The "universal" priming sequences used in the examples presented infra were each 16 nt long.

The genomic DNA to be used as substrate for amplification will come from the eukaryotic species from which the genomic sequence data had originally been

obtained, or a closely related species, and can conveniently be prepared by well known techniques from somatic or germline tissue or cultured cells of the organism. See, e.g., Short Protocols in Molecular Biology: A Compendium of Methods from Current Protocols in Molecular Biology, Ausubel et al. (eds.), 4th edition (April 1999), John Wiley & Sons (ISBN: 047132938X) and Maniatis et al., Molecular Cloning: A Laboratory Manual, 2nd edition (December 1989), Cold Spring Harbor Laboratory Press (ISBN: 0879693096). Many such prepared genomic DNAs are available commercially, with the human genomic DNAs additionally having certification of donor informed consent.

Although the intronic and intergenic material

flanking putative coding regions in the amplicons could

potentially interfere with hybridizations during microarray
experiments, we have found, surprisingly, that differential
expression ratios are not significantly affected. Rather,
the predominant effect of exon size is to alter the

absolute signal intensity, rather than its ratio. Equally
surprising, the art had suggested that single exon probes
would not provide sufficient signal intensity for high
stringency hybridization analyses; we find that such probes
not only provide adequate signal, but have substantial

advantages, as herein described.

After partial purification, as by size exclusion spin column, with or without confirmation as to amplicon quality as by gel electrophoresis, each amplicon (single exon probe) is disposed in an array upon a support substrate.

Methods for creating microarrays by deposition and fixation of nucleic acids onto support substrates are well known in the art (Reviewed by Schena et al., see above).

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Typically, the support substrate will be glass,

although other materials, such as amorphous or crystalline silicon or plastics. Such plastics include polymethylacrylic, polyethylene, polypropylene, polyacrylate, polymethylmethacrylate, polyvinylchloride, polytetrafluoroethylene, polystyrene, polycarbonate, polyacetal, polysulfone, celluloseacetate, cellulosenitrate, nitrocellulose, or mixtures thereof, can also be used. Typically, the support will be rectangular, although other shapes, particularly circular disks and even spheres, present certain advantages. Particularly advantageous alternatives to glass slides as support substrates for array of nucleic acids are optical discs, as described in WO 98/12559.

The amplified nucleic acids can be attached

15 covalently to a surface of the support substrate or, more
typically, applied to a derivatized surface in a chaotropic
agent that facilitates denaturation and adherence by
presumed noncovalent interactions, or some combination
thereof.

Robotic spotting devices useful for arraying nucleic acids on support substrates can be constructed using public domain specifications (The MGuide, version 2.0, http://cmgm.stanford.edu/pbrown/mguide/index.html), or can conveniently be purchased from commercial sources

(MicroArray GenII Spotter and MicroArray GenIII Spotter, Molecular Dynamics, Inc., Sunnyvale, CA). Spotting can also be effected by printing methods, including those using ink jet technology.

As is well known in the art, microarrays

typically also contain immobilized control nucleic acids.

For controls useful in providing measurements of background signal for the genome-derived single exon microarrays of the present invention, a plurality of *E. coli* genes can readily be used. As further described in Example 1, 16 or

32 *E. coli* genes suffice to provide a robust measure of

background noise in such microarrays.

As is well known in the art, the amplified product disposed in arrays on a support substrate to create a nucleic acid microarray can consist entirely of natural nucleotides linked by phosphodiester bonds, or alternatively can include either nonnative nucleotides, alternative internucleotide linkages, or both, so long as complementary binding can be obtained in the hybridization. If enzymatic amplification is used to produce the immobilized probes, the amplifying enzyme will impose certain further constraints upon the types of nucleic acid analogs that can be generated.

Although particularly described herein as using high density microarrays constructed on planar substrates, the methods of the present invention for confirming the expression of ORFs predicted from genomic sequence can use any of the known types of microarrays, as herein defined, including lower density planar arrays, and microarrays on nonplanar, nonunitary, distributed substrates.

20 For example, gene expression can be confirmed using hybridization to lower density arrays, such as those constructed on membranes, such as nitrocellulose, nylon, and positively-charged derivatized nylon membranes.

Further, gene expression can also be confirmed using nonplanar, bead-based microarrays such as are described in Brenner et al., Proc. Natl. Acad. Sci. USA 97(4):166501670 (2000); U.S. Patent No. 6,057,107; and U.S. Patent No. 5,736,330. In theory, a packed collection of such beads provides in aggregate a higher density of nucleic acid probe than can be achieved with spotting or lithography techniques on a single planar substrate.

Planar microarrays on solid substrates, however, provide certain useful advantages, including high throughput and compatibility with existing readers. For example, each standard microscope slide can include at

least 1000, typically at least 2000, preferably 5000 and upto 10,000 - 50,000 or more nucleic acid probes of discrete sequence. The number of sequences deposited will depend on their required application.

Each putative gene can be represented in the array by a single predicted ORF. Alternatively, genes can be represented by more than one predicted ORF. For purposes of measuring differential splicing, more than one predicted ORF will be provided for a putative gene. And as is well known in the art, each probe of defined sequence, representing a single predicted ORF, can be deposited in a plurality of locations on a single microarray to provide redundancy of signal.

The genome-derived single exon microarrays

described above differ in several fundamental and
advantageous ways from microarrays presently used in the
gene expression art, including (1) those created by
deposition of mRNA-derived nucleic acids, (2) those created
by in situ synthesis of oligonucleotide probes, and (3)

those constructed from yeast genomic DNA.

Most nucleic acid microarrays that are in use for study of eukaryotic gene expression have as immobilized probes nucleic acids that are derived — either directly or indirectly — from expressed message. As discussed above, it is common, for example, for such microarrays to be derived from cDNA/EST libraries, either from those previously described in the literature, see Lennon et al., or from the de novo construction of "problem specific" libraries targeted at a particular biological question, R.S. Thomas et al., Cancer Res. (in press). Such microarrays are herein collectively denominated "EST microarrays".

Such EST microarrays by definition can measure expression only of those genes found in EST libraries,

35 shown herein to represent only a fraction of expressed

genes. Furthermore, such libraries — and thus microarrays based thereupon — are biased by the tissue or cell type of message origin, by the expression levels of the respective genes within the tissues, and by the ability of the message successfully to have been reverse-transcribed and cloned.

Thus, as further discussed in Example 1, the methods of the present invention enable sequences that do not appear in EST or other expression databases to be determined - subsequently arrayed for expression

10 measurements could not, therefore, have been represented as probes on an EST microarray. And as further demonstrated in the examples, infra, the remaining population of genes identified from genomic sequence by the methods of the present invention — that is, the one third of sequences that had previously been accessioned in EST or other expression databases — are biased toward genes with higher expression levels.

Representation of a message in an EST and/or cDNA library depends upon the successful reverse transcription, optionally but typically with subsequent successful cloning, of the message. This introduces substantial bias into the population of probes available for arraying in EST—microarrays.

In contrast, neither reverse transcription nor

cloning is required to produce the probes arrayed on the
genome-derived single exon microarrays of the present
invention. And although the ultimate deposition of a probe
on the genome-derived single exon microarray of the present
invention depends upon a successful amplification from

genomic material, a priori knowledge of the sequence of the
desired amplicon affords greater opportunity to recover any
given probe sequence recalcitrant to amplification than is
afforded by the requirement for successful reverse
transcription and cloning of unknown message in EST
approaches.

Thus, the genome-derived single exon microarrays of the present invention present a far greater diversity of probes for measuring gene expression, with far less bias, than do EST microarrays presently used in the art.

As a further consequence of their ultimate origin from expressed message, the probes in EST microarrays often contain poly-A (or complementary poly-T) stretches derived from the poly-A tail of mature mRNA. These homopolymeric stretches contribute to cross-hybridization, that is, to a spurious signal occasioned by hybridization to the homopolymeric tail of a labeled cDNA that lacks sequence homology to the gene-specific portion of the probe.

In contrast, the probes arrayed in the genomederived single exon microarrays of the present invention

15 lack homopolymeric stretches derived from message
polyadenylation, and thus can provide more specific signal.

Typically, at least about 50, 60 or 75% of the probes on
the genome-derived single exon microarrays of the present
invention lack homopolymeric regions consisting of A or T,

20 where a homopolymeric region is defined for purposes herein
as stretches of 25 or more, typically 30 or more, identical
nucleotides.

A further distinction, which also affects the specificity of hybridization, is occasioned by the typical derivation of EST microarray probes from cloned material. Because much of the probe material disposed as probes on EST microarrays is excised or amplified from plasmid, phage, or phagemid vectors, EST microarrays typically include a fair amount of vector sequence, more so when the probes are amplified, rather than excised, from the vector.

In contrast, the vast majority of probes in the genome-derived single exon microarrays of the present invention contain no prokaryotic or bacteriophage vector sequence, having been amplified directly or indirectly from genomic DNA. Typically, therefore, at least about 50, 60,

70 or 80% or more of individual exon-including probes

disposed on a genome-derived single exon microarray of the present invention lack vector sequence, and particularly lack sequences drawn from plasmids and bacteriophage. 5 Preferably, at least about 85, 90 or more than 90% of exonincluding probes in the genome-derived single exon microarray of the present invention lack vector sequence. With attention to removal of vector sequences through preprocessing 24, percentages of vector-free exon-including 10 probes can be as high as 95 - 99%. The substantial absence of vector sequence from the genome-derived single exon microarrays of the present invention results in greater specificity during hybridization, since spurious crosshybridization to a probe vector sequence is reduced.

As a further consequence of excision or amplification of probes from vectors in construction of EST microarrays, the probes arrayed thereon often contain artificial sequence, derived from vector polylinker multiple cloning sites, at both 5' and 3' ends. The probes 20 disposed upon the genome-derived single exon microarrays need have no such artificial sequence appended thereto.

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As mentioned above, however, the ORF-specific primers-used-to-amplify-putative-ORFs-can-includeartificial sequences, typically 5' to the ORF-specific 25 primer sequence, useful for "universal" (that is, independent of ORF sequence) priming of subsequent amplification or sequencing reactions. When such "universal" 5' and/or 3' priming sequences are appended to the amplification primers, the probes disposed upon the 30 genome-derived single exon microarray will include artificial sequence similar to that found in EST microarrays. However, the genome-derived single exon microarray of the present invention can be made without such sequences, and if so constructed, presents an even 35 smaller amount of nonspecific sequence that would

contribute to nonspecific hybridization.

Yet another consequence of typical use of cloned material as probes in EST microarrays is that such microarrays contain probes that result from cloning

5 artifacts, such as chimeric molecules containing coding region of two separate genes. Derived from genomic material, typically not thereafter cloned, the probes of the genome-derived single exon microarrays of the present invention lack such cloning artifacts, and thus provide greater specificity of signal in gene expression measurements.

A further consequence of the cloned origin of probes on many EST microarrays is that the individual probes often have disparate sizes, which can cause the optimal hybridization stringency to vary among probes on a single microarray. In contrast, as discussed above, the probes arrayed on the genome-derived single exon microarrays of the present invention can readily be designed to have a narrow distribution in sizes, with the range of probe sizes no greater than about 10% of the average size, typically no greater than about 5% of the average probe size.

Because of their origin from fully- or partiallyspliced message, probes disposed upon EST arrays will often
include multiple exons. The percentage of such exonspanning probes in an EST microarray can be calculated, on
average, based upon the predicted number of exons/gene for
the given species and the average length of the immobilized
probes. For human genes, the near-complete sequence of
human chromosome 22, Dunham et al., Nature 402(6761):489-95
(1999), predicts that human genes average 5.5 exons/gene.
Even with probes of 200 - 500 bp, the vast majority of
human EST microarray probes include more than one exon.

In contrast, by virtue of their origin from algorithmically identified ORFs in genomic sequence, the

probes in the genome-derived single exon microarrays of the present invention can consist of individual exons. Thus, in contrast to EST microarrays, at least about 50, 60, 70, 75, 80, 85, 95 or 99% of probes deposited in the genome-5 derived microarray of the present invention consist of, or include, no more than one predicted ORF.

This provides the ability, not readily achieved using EST microarrays, to use the genome-derived single exon microarrays of the present invention to measure 10 tissue-specific expression of individual exons, which in turn allows differential splicing events to be detected and characterized, and in particular, allows the correlation of differential splicing to tissue-specific expression patterns.

Furthermore, the exons that are represented in EST microarrays are often biased toward the 3' or 5' end of their respective genes, since sequencing strategies used for EST identification are so biased. In contrast, no such 3' or 5' bias necessarily inheres in the selection of exons 20 for disposition on the genome-derived single exon microarrays of the present invention.

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Conversely, the probes provided on the genomederived single exon microarrays of the present invention typically, but need not necessarily, include intronic 25 and/or intergenic sequence that is absent from EST microarrays, which are derived from mature mRNA. Typically, at least about 50, 60, 70, 80 or 90% of the exon-including probes on the genome-derived single exon microarrays of the present invention include sequence drawn 30 from noncoding regions. As discussed above, the additional presence of noncoding region does not significantly interfere with measurement of gene expression, and provides the additional opportunity to assay prespliced RNA, and thus measure such phenomena such as nuclear export control.

The genome-derived single exon microarrays of the

present invention are also quite different from in situ synthesis microarrays, where probe size is severely constrained by inadequacies in the photolithographic synthesis process.

Typically, probes arrayed on in situ synthesis microarrays are limited to a maximum of about 25 bp. As a well known consequence, hybridization to such chips must be performed at low stringency. In order, therefore, to achieve unambiguous sequence-specific hybridization results, the in situ synthesis microarray requires substantial redundancy, with concomitant programmed arraying for each probe of probe analogues with altered (i.e., mismatched) sequence.

In contrast, the longer probe length of the
genome-derived single exon microarrays of the present
invention allows much higher stringency hybridization and
wash. Typically, therefore, exon-including probes on the
genome-derived single exon microarrays of the present
invention average at least about 100, 200, 300, 400 or
500 bp in length. By obviating the need for substantial
probe redundancy, this approach permits a higher density of
probes for discrete exons or genes to be arrayed on the
microarrays of the present invention than can be achieved
for in situ synthesis microarrays.

A further distinction is that the probes in in situ synthesis microarrays typically are covalently linked to the substrate surface. In contrast, the probes disposed on the genome-derived microarray of the present invention typically are, but need not necessarily be, bound noncovalently to the substrate.

Furthermore, the short probe size on in situ microarrays causes large percentage differences in the melting temperature of probes hybridized to their complementary target sequence, and thus causes large percentage differences in the theoretically optimum

stringency across the array as a whole.

In contrast, the larger probe size in the microarrays of the present invention create lower percentage differences in melting temperature across the range of arrayed probes.

A further significant advantage of the microarrays of the present invention over in situ synthesized arrays is that the quality of each individual probe can be confirmed before deposition. In contrast, the quality of probes cannot be assessed on a probe-by-probe basis for the in situ synthesized microarrays presently being used.

The genome-derived single exon microarrays of the present invention are also distinguished over, and present substantial benefits over, the genome-derived microarrays from lower eukaryotes such as yeast. Lashkari et al., Proc. Natl. Acad. Sci. USA 94:13057-13062 (1997).

Only about 220 - 250 of the 6100 or so nuclear genes in Saccharomyces cerevisiae — that is, only about 4

20 - 5% — have standard, spliceosomal, introns, Lopez et al.,

Nucl. Acids Res. 28:85-86 (2000); Spingola et al., RNA

5(2):221-34 (1999). Furthermore, the entire yeast genome

has already been sequenced. These two facts permit the

ready amplification and disposition of single-ORF amplicons

25 on such microarray without the requirement for antecedent

use of gene prediction and/or comparative sequence

analyses.

Thus, a significant aspect of the present invention is the ability to identify and to confirm

30 expression of predicted coding regions in genomic sequence drawn from eukaryotic organisms that have a higher percentage of genes having introns than do yeast such as Saccharomyces cerevisiae, particularly in genomic sequence drawn from eukaryotes in which at least about 10, 20 or 50% of protein-encoding genes have introns. In preferred

embodiments, the methods and apparatus of the present invention are used to identify and confirm expression of novel genes from genomic sequence of eukaryotes in which the average number of introns per gene is at least about one, two or three or more.

After the physical substrate is prepared, experimental verification of predicted function is performed.

In a preferred embodiment of the present
invention, where the function sought to be identified in
genomic sequence is protein coding, experimental
verification is performed by measuring expression of the
putative ORFs, typically through nucleic acid hybridization
experiments, and in particularly preferred embodiments,
through hybridization to genome-derived single exon
microarrays prepared as above- described.

for each probe in the microarray as a ratio of the expression measured concurrently in a plurality of mRNA

20 sources, according to techniques well known in the microarray art, Reviewed in Schena et al., and as further described in Example 2, below. The mRNA source for the reference against which specific expression is measured can be drawn from a homogeneous mRNA source, such as a single cultured cell-type, or alternatively can be heterogeneous, as from a pool of mRNA derived from multiple tissues and/or cell types, as further described in Example 2, infra.

mRNA can be prepared by standard techniques, see
Ausubel et al. and Maniatis et al., or purchased

commercially. The mRNA is then typically reversetranscribed in the presence of labeled nucleotides: the
index source (that in which expression is desired to be
measured) is reverse transcribed in the presence of
nucleotides labeled with a first label, typically a

fluorophore (fluorochrome; fluor; fluorescent dye); the

reference source is reverse transcribed in the presence of a second label, typically a fluorophore, typically fluorometrically-distinguishable from the first label. As further described in Example 2, infra, Cy3 and Cy5 dyes 5 prove particularly useful in these methods. After partial purification of the index and reference targets, hybridization to the probe array is conducted according to standard techniques, typically under a coverslip.

After wash, microarrays are conveniently scanned 10 using a commercial microarray scanning device, such as a Gen3 Scanner (Molecular Dynamics, Sunnyvale, CA). Data on expression is then passed, with or without interim storage, to process 500, where the results for each probe are related to the original sequence.

15

Often, hybridization of target material to the genome-derived single exon microarray will identify certain of the probes thereon as of particular interest. Thus, it is often desirable that the user be able readily to obtain sufficient quantities of an individual probe, either for 20 subsequent arrayed deposition upon an additional support substrate, often as part of a microarray having a plurality of probes so identified, or alternatively or additionally as a solitary solid-phase or solution-phase probe, for further use.

25 Thus, in another aspect, the present invention provides compositions and kits for the ready production of nucleic acids identical in sequence to, or substantially identical in sequence to, probes on the genome-derived single exon microarrays of the present invention.

30 In this aspect, a small quantity of each probe is disposed, typically without attachment to substrate, in a spatially-addressable ordered set, typically one per well of a microtiter dish. Although a 96 well microtiter plate can be used, greater efficiency is obtained using higher 35 density arrays, such as are provided by microtiter plates

having 384, 864, 1536, 3456, 6144, or 9600 wells, and although microtiter plates having physical depressions (wells) are conveniently used, any device that permits addressable withdrawal of reagent from fluidly-noncommunicating areas can be used.

In this aspect of the invention, therefore, a fluidly noncommunicating addressable ordered set of individual probes, corresponding to those on a genomederived single exon microarray, is provided, with each probe in sufficient quantity to permit amplification, such as by PCR. As earlier mentioned, the ORF-specific 5' primers used for genomic amplification can have a first common sequence added thereto, and the ORF-specific 3' primers used for genomic amplification can have a second, different, common sequence added thereto, thus permitting, in this preferred embodiment, the use of a single set of 5' and 3' primers to amplify any one of the probes from the amplifiable ordered set.

Each discrete amplifiable probe can also be

20 packaged with amplification primers, solutes, buffers,
etc., and can be provided in dry (e.g., lyophilized) form
or wet, in the latter case typically with addition of
agents that retard evaporation.

In another aspect of the present invention, a

25 genome-derived single-exon microarray is packaged together
with such an ordered set of amplifiable probes
corresponding to the probes, or one or more subsets of
probes, thereon. In alternative embodiments, the ordered
set of amplifiable probes is packaged separately from the
30 genome-derived single exon microarray.

In some embodiments, the microarray and/or ordered probe set are further packaged with recordable media that provide probe identification and addressing information, and that can additionally contain annotation information, such as gene expression data. Such recordable

media can be packaged with the microarray, with the ordered probe set, or with both.

If the microarray is constructed on a substrate that incorporates recordable media, such as is described in international patent application no. WO 98/12559, then separate packaging of the genome-derived single exon microarray and the bioinformatic information is not required.

The amount of amplifiable probe material should to be sufficient to permit at least one amplification sufficient for subsequent hybridization assay.

Although the use of high density genome-derived microarrays on solid planar substrates is presently a preferred approach for the physical confirmation and characterization of the expression of sequences predicted to encode protein, other types of microarrays (as herein defined) can also be used.

Furthermore, as earlier mentioned, experimental verification of the function predicted from genomic sequence in process 200 can be bioinformatic, rather than, or additional to, physical verification.

For example, where the function desired to be identified is protein coding, the predicted ORFs can be compared bioinformatically to sequences known or suspected of being expressed.

Thus, the sequences output from process 300 (or process 200), can be used to query expression databases, such as EST databases, SNP ("single nucleotide polymorphism") databases, known cDNA and mRNA sequences, SAGE ("serial analysis of gene expression") databases, and more generalized sequence databases that allow query for expressed sequences. Such query can be done by any sequence query algorithm, such as BLAST ("basic local alignment search tool"). The results of such query — including information on identical sequences and

information on nonidentical sequences that have diffuse or focal regions of sequence homology to the query sequence can then be passed directly to process 500, or used to inform analyses subsequently undertaken in process 200, 5 process 300, or process 400.

Experimental data, whether obtained by physical or bioinformatic assay in process 400, is passed to process 500 where it is usefully related to the sequence data itself, a process colloquially termed "annotation". Such 10 annotation can be done using any technique that usefully relates the functional information to the sequence, as, for example, by incorporating the functional data into the record itself, by linking records in a hierarchical or relational database, by linking to external databases, or 15 by a combination thereof. Such database techniques are well within the skill in the art.

The annotated sequence data can be stored locally, uploaded to genomic sequence database 100, and/or displayed 800.

The methods and apparatus of the present invention rapidly produce functional information from genomic sequence. Coupled with the escalating pace at which sequence now accumulates, the rapid pace of sequence annotation produces a need for methods of displaying the 25 information in meaningful ways.

20

FIG. 3 shows visual display 80 presenting a single genomic sequence annotated according to the present Because of its nominal resemblance to artistic works of Piet Mondrian, visual display 80 is alternatively 30 described herein as a "Mondrian".

Each of the visual elements of display 80 is aligned with respect to the genomic sequence being annotated (hereinafter, the "annotated sequence"). Given the number of nucleotides typically represented in an annotated sequence, representation of individual

nucleotides would rarely be readable in hard copy output of display 80. Typically, therefore, the annotated sequence is schematized as rectangle 89, extending from the left border of display 80 to its right border. By convention herein, the left border of rectangle 89 represents the first nucleotide of the sequence and the right border of rectangle 89 represents the last nucleotide of the sequence.

As further discussed below, however, the Mondrian visual display of annotated sequence can serve as a convenient graphical user interface for computerized representation, analysis, and query of information stored electronically. For such use, the individual nucleotides can conveniently be linked to the X axis coordinate of rectangle 89. This permits the annotated sequence at any point within rectangle 89 readily to be viewed, either automatically — for example, by time-delayed appearance of a small overlaid window upon movement of a cursor or other pointer over rectangle 89 — or through user intervention, as by clicking a mouse or other pointing device at a point in rectangle 89.

Visual display 80 is generated after user

specification of the genomic sequence to be displayed.

Such specification can consist of or include an accession

25 number for a single clone (e.g., a single BAC accessioned into GenBank), wherein the starting and stopping nucleotides are thus absolutely identified, or alternatively can consist of or include an anchor or fulcrum point about which a chosen range of sequence is

30 anchored, thus providing relative endpoints for the sequence to be displayed. For example, the user can anchor such a range about a given chromosomal map location, gene name, or even a sequence returned by query for similarity or identity to an input query sequence. When visual

35 display 80 is used as a graphical user interface to

computerized data, additional control over the first and last displayed nucleotide will typically be dynamically selectable, as by use of standard zooming and/or selection tools.

5 Field 81 of visual display 80 is used to present the output from process 200, that is, to present the bioinformatic prediction of those sequences having the desired function within the genomic sequence. Functional sequences are typically indicated by at least one rectangle 83 (83a, 83b, 83c), the left and right borders of which respectively indicate, by their X-axis coordinates, the starting and ending nucleotides of the region predicted to have function.

Where a single bioinformatic method or approach
identifies a plurality of regions having the desired
function, a plurality of rectangles 83 is disposed
horizontally in field 81. Where multiple methods and/or
approaches are used to identify function, each such method
and/or approach can be represented by its own series of:
horizontally disposed rectangles 83, each such horizontally
disposed series of rectangles offset vertically from those
representing the results of the other methods and
approaches.

Thus, rectangles 83a in FIG. 3 represent the

25 functional predictions of a first method of a first
approach for predicting function, rectangles 83b represent
the functional predictions of a second method and/or second
approach for predicting that function, and rectangles 83c
represent the predictions of a third method and/or

30 approach.

Where the function desired to be identified is protein coding, field 81 is used to present the bioinformatic prediction of sequences encoding protein.

For example, rectangles 83a can represent the results from GRAIL or GRAIL II, rectangles 83b can represent the results

from GENEFINDER, and rectangles 83c can represent the results from DICTION.

Optionally, and preferably, rectangles 83 collectively representing predictions of a single method and/or approach are identically colored and/or textured, and are distinguishable from the color and/or texture used for a different method and/or approach.

Alternatively, or in addition, the color, hue, density, or texture of rectangles 83 can be used further to report a measure of the bioinformatic reliability of the prediction. For example, many gene prediction programs will report a measure of the reliability of prediction. Thus, increasing degrees of such reliability can be indicated, e.g., by increasing density of shading. Where display 80 is used as a graphical user interface, such measures of reliability, and indeed all other results output by the program, can additionally or alternatively be made accessible through linkage from individual rectangles 83, as by time-delayed window ("tool tip" window), or by pointer (e.g., mouse)-activated link.

As earlier described, increased predictive reliability can be achieved by requiring consensus among methods and/or approaches to determining function. Thus, field 81 can include a horizontal series of rectangles 83 that indicate one or more degrees of consensus in predictions of function.

Although FIG. 3 shows three series of horizontally disposed rectangles in field 81, display 80 can include as few as one such series of rectangles and as 30 many as can discriminably be displayed, depending upon the number of methods and/or approaches used to predict a given function.

Furthermore, field 81 can be used to show predictions of a plurality of different functions.

35 However, the increased visual complexity occasioned by such

display makes more useful the ability of the user to select a single function for display. When display 80 is used as a graphical user interface for computer query and analysis, such function can usefully be indicated and user-5 selectable, as by a series of graphical buttons or tabs (not shown in FIG. 3).

Rectangle 89 is shown in FIG. 3 as including interposed rectangle 84. Rectangle 84 represents the portion of annotated sequence for which predicted 10 functional information has been assayed physically, with the starting and ending nucleotides of the assayed material indicated by the X axis coordinates of the left and right borders of rectangle 84. Rectangle 85, with optional inclusive circles 86 (86a, 86b, and 86c) displays the 15 results of such physical assay.

Although a single rectangle 84 is shown in FIG. 3, physical assay is not limited to just one region of annotated genomic sequence. It is expected that an increasing percentage of regions predicted to have function 20 by process 200 will be assayed physically, and that display 80 will accordingly, for any given genomic sequence, have an increasing number of rectangles 84 and 85, representing an-increased-density-of-sequence-annotation.---

Where the function desired to be identified is 25 protein coding, rectangle 84 identifies the sequence of the probe used to measure expression. In embodiments of the present invention where expression is measured using genome-derived single exon microarrays, rectangle 84 identifies the sequence included within the probe 30 immobilized on the support surface of the microarray. As noted supra, such probe will often include a small amount of additional, synthetic, material incorporated during amplification and designed to permit reamplification of the probe, which sequence is typically not shown in display 80. 35

bioinformatic assay of the genomic sequence. For example, where the function desired to be identified is protein coding, process 400 can include bioinformatic query of expression databases with the sequences predicted in process 200 to encode exons. And as earlier discussed, because bioinformatic assay presents fewer constraints than does physical assay, often the entire output of process 200 can be used for such assay, without further subsetting thereof by process 300. Therefore, rectangle 87 typically need not have separate indicators therein of regions submitted for bioinformatic assay; that is, rectangle 87 typically need not have regions therein analogous to rectangles 84 within rectangle 89.

Rectangle 87 as shown in FIG. 3 includes smaller
rectangles 880 and 88. Rectangles 880 indicate regions
that returned a positive result in the bioinformatic assay,
with rectangles 88 representing regions that did not return
such positive results. Where the function desired to be
predicted and displayed is protein coding, rectangles 880
indicate regions of the predicted exons that identify
sequence with significant similarity in expression
databases, such as EST, SNP, SAGE databases, with
rectangles 88 indicating genes novel over those identified
in existing expression data bases.

Rectangles 880 can further indicate, through color, shading, texture, or the like, additional information obtained from bioinformatic assay.

25

For example, where the function assayed and displayed is protein coding, the degree of shading of rectangles 880 can be used to represent the degree of sequence similarity found upon query of expression databases. The number of levels of discrimination can be as few as two (identity, and similarity, where similarity has a user-selectable lower threshold). Alternatively, as many different levels of discrimination can be indicated as

can visually be discriminated.

Where display 80 is used as a graphical user interface, rectangles 880 can additionally provide links directly to the sequences identified by the query of expression databases, and/or statistical summaries thereof. As with each of the precedingly-discussed uses of display 80 as a graphical user interface, it should be understood that the information accessed via display 80 need not be resident on the computer presenting such display, which often will be serving as a client, with the linked information resident on one or more remotely located servers.

Rectangle 85 displays the results of physical assay of the sequence delimited by its left and right borders.

Rectangle 85 can consist of a single rectangle, thus indicating a single assay, or alternatively, and increasingly typically, will consist of a series of rectangles (85a, 85b, 85c) indicating separate physical assays of the same sequence.

Where the function assayed is gene expression, and where gene expression is assayed as herein described using simultaneous two-color fluorescent detection of hybridization to genome-derived single exon microarrays, individual rectangles 85 can be colored to indicate the degree of expression relative to control. Conveniently, shades of green can be used to depict expression in the sample over control values, and shades of red used to depict expression less than control, corresponding to the spectra of the Cy3 and Cy5 dyes conventionally used for respective labeling thereof. Additional functional information can be provided in the form of circles 86 (86a, 86b, 86c), where the diameter of the circle can be used to indicate expression intensity. As discussed infra, such relative expression (expression ratios) and absolute

expression (signal intensity) can be expressed using normalized values.

where display 80 is used as a graphical user interface, rectangle 85 can be used as a link to further information about the assay. For example, where the assay is one for gene expression, each rectangle 85 can be used to link to information about the source of the hybridized mRNA, the identity of the control, raw or processed data from the microarray scan, or the like.

FIG. 4 is rendition of display 80 representing 10 gene prediction and gene expression for a hypothetical BAC, showing conventions used in the Examples presented infra. BAC sequence ("Chip seq.") 89 is presented, with the physically assayed region thereof (corresponding to rectangle 84 in FIG. 3) shown in white. Algorithmic gene predictions are shown in field 81, with predictions by GRAIL shown, predictions by GENEFINDER, and predictions by DICTION shown. Within rectangle 87, regions of sequence that, when used to query expression databases, return 20 identical or similar sequences ("EST hit") are shown as white rectangles (corresponding to rectangles 880 in FIG. 3), gray indicates low homology, and black indicates unknowns (where black and gray would correspond to rectangles 88 in FIG. 3).

Although FIGS. 3 and 4 show a single stretch of sequence, uninterrupted from left to right, longer sequences are usefully represented by vertical stacking of such individual Mondrians, as shown in FIGS. 9 and 10.

## 30 Single Exon Probes Useful For Measuring Gene Expression

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The methods and apparatus of the present invention rapidly produce functional information from genomic sequence. Where the function to be identified is protein coding, the methods and apparatus of the present

invention rapidly identify and confirm the expression of portions of genomic sequence that function to encode protein. As a direct result, the methods and apparatus of the present invention rapidly yield large numbers of single-exon nucleic acid probes, the majority from previously unknown genes, each of which is useful for measuring and/or surveying expression of a specific gene in one or more tissues or cell types.

It is, therefore, another aspect of the present invention to provide genome-derived single exon nucleic acid probes useful for gene expression analysis, and particularly for gene expression analysis by microarray.

Using the methods and genome-derived single-exon microarrays of the present invention, we have for example readily identified a large number of unique ORFs from human genomic sequence. Using single exon probes that encompass these ORFs, we have demonstrated, through microarray hybridization analysis, the expression of 5,074 of these ORFs in HBL 100 cells.

20 The HBL 100 human breast cancer cell line was established in vitro from milk of an apparently healthy woman. The cells express a variant of SV40 large T antigen, and genomic DNA from HBL-100 cells possesses transforming activity associated with the viral 25 information. The HBL100 cell line is nontumorigenic, and acquires the capacity to invade normal tissues and to replace them by proliferation in vitro only at high passage levels (HPL); these epithelial cells are thus are a useful model for studying breast tumor progression in vitro. HBL 100 cells bind both epidermal growth factor (EGF) and qlucocorticoids but are progesterone receptor negative.

As would immediately be appreciated by one of skill in the art, each single exon probe having demonstrable expression in HBL 100 cells is currently available for use in measuring the level of its ORF's expression in Breast.

Diseases of the breast are a significant cause of human morbidity and mortality. Increasingly, genetic factors are being found that contribute to predisposition, onset, and/or aggressiveness of most, if not all, of these diseases. Although mutations in single genes have been identified as causative for some diseases of the breast, for the most part these disorders are believed to have polygenic etiologies.

For example, according to the American Cancer

10 Society (ACS), carcinoma of the breast is the second most common cancer in women and, after lung cancer, is the second deadliest. The ACS estimates that in the U.S. there occurred 182,800 new cases of malignant breast cancer in 2000, and about 40,800 deaths from the disease. Although incidence of breast cancer is said to have declined, the disease clearly continues to represent a serious risk to the health and life of American women. Indeed, about one in nine U.S. women will develop breast cancer in her lifetime, and at present mortality rates, about a third of such women will eventually die from the disease.

A variety of factors are known to increase the risk of breast carcinoma. Sex is one: breast cancer in men is rare. Age is another: as women age, their risk for developing breast cancer increases, a 70 year old woman having three times the risk of developing cancer and five times the risk of dying from the disease as compared to a 40 year old woman. Most breast cancers occur after age 50, although in women with a genetic susceptibility, breast cancer tends to occur at an earlier age than in sporadic cases. Reproductive and menstrual history are also known to affect risk, with risk increasing with early menarche and late menopause, and is reduced by early first full term pregnancy. Additional risk factors, oft-times termed "lifestyle factors", include weight gain, obesity, fat intake, alcohol consumption, and level of physical

activity.

That genetic factors underlie the etiology of breast cancer is suggested by the approximately two-fold increased risk for development of breast cancer by women with a first-degree relative who has also developed breast cancer. After gender and age, a positive family history is the strongest known predictive risk factor for breast cancer. Genetic linkage analysis in families with high rates of inherited cancer have facilitated the identification of several genes in which mutations can be shown to contribute substantially to the development and progression of breast cancer, including BRCA1, BRCA2, p53, and PTEN/MMAC1. Further study has made clear, however, that these genes are not alone sufficient to explain all genetic contributions to breast cancer.

For example, BRCA1 appears to be responsible for disease in up to 90% of families with both breast and ovarian cancer, but in only 45% of families with multiple cases of breast cancer without occurrence of ovarian

20 cancer. And mutations in BRCA2, localized to the long arm of chromosome 13, are thought to account for only approximately 35% of multiple case breast cancer families.

Furthermore, despite the strong correlation between germline mutations in BRCA1 or BRCA2 and development of

25 breast cancer, only weak connections have been made between these genes and sporadic breast cancer.

Epistatic effects of BRCA1 and BRCA2 mutations on other genetic loci, only some of which have been identified, have been postulated to account for some of the deleterious effects of mutations in these two genes.

Thus, mutations in p53 seem to be much more frequent in BRCA1 breast cancers (20/26) and somewhat more frequent in BRCA2-associated breast cancers (10/22) than in grade-matched sporadic cancers (7/20). BRCA mutation-associated cancers contain p53 mutations not typically

found in sporadic breast cancer, and 12 individual hereditary breast cancers have been shown to contain more than a single p53 mutation. Mutations of BRCA1 and BRCA2 may thus confer a "mutator" phenotype permitting the accumulation of genetic abnormalities, with p53 inactivation selected during tumor progression.

Additionally, genome-wide screening for chromosomal gains or losses in breast cancers harboring BRCA1 or BRCA2 mutations demonstrated more regions that were amplified or deleted compared to controls, suggesting a generalized increase in large-scale genomic instability. Chromosomes 5q, 4q, and 4p had very frequent loss of heterozygosity in BRCA1 tumors, while BRCA2 tumors were characterized by losses at 13q (near the BRCA2 locus itself) and 6q, and chromosomal gains at 17q (outside of the HER2/neu locus) and 20q.

Mutations of other genes have also been implicated in susceptibility to development or aggressiveness of breast cancer. For example, germline mutations in the ATM gene, localized to chromosome 11q22-23, result in an increased risk of breast cancer among female heterozygote carriers with an estimated relative-risk-of-3.9-to-6.4; it-is-unclear, however, if-mutations in the ATM gene itself contribute to breast cancer.

Normal allelic variation in a variety of genes, as opposed to frank mutation, may also influence susceptibility to developing breast carcinoma and the propensity for the disease to progress. Such 30 polymorphisms may thus explain why particular women or ethnic groups who do not otherwise bear mutations in genes known to be linked to breast cancer are at greater risk, especially in the context of exposure to environmental agents and other nonhereditary risk factors.

35 Polymorphically expressed genes may code for enzymes that

metabolize estrogens or detoxify drugs and environmental carcinogens.

For example, molecular epidemiologic studies of cancer of the breast have examined associations with p450 cytochrome genotypes including CYP1A1, CYP2D6, and CYP17. The CYP1A1 gene, located on chromosome 15q, encodes the enzyme aryl hydrocarbon hydroxylase (AHH), present in breast tissue, and which catabolizes polycyclic aromatic hydrocarbons and arylamines. AHH is strongly inducible, i.e., greater enzymatic activity is seen with greater exposure to substrates. AHH catalyzes the monooxygenation of polycyclic aromatic hydrocarbons to phenolic products and epoxides that may be carcinogenic. AHH is also involved in the conversion of estrogen to hydroxylated conjugated estrogens such as 2-hydroxyestradiol.

Three polymorphisms in the CYP1A1 gene have been identified: an MspI RFLP of the 3' end of the gene (MspI); an adenine to guanine mutation in exon 7, causing an isoleucine to valine substitution (Ile-Val); and a polymorphism of the CYP1A1 gene identified among Negroids. The frequencies of the MspI and Ile-Val polymorphisms vary considerably by race, being higher among Japanese and Hawaiian populations as compared with Caucasians and Negroids.

25 The CYP2D6 gene is located on chromosome 22q and encodes the enzyme debrisoquine hydroxylase, which metabolizes a variety of drugs and other xenobiotics. Like other polymorphically expressed p450 enzymes, it may activate procarcinogens or, conversely, detoxify

30 carcinogens. A number of alleles have been characterized at the CYP2D6 locus. The "poor metabolizer" phenotype (CYP2D6 mutant/mutant genotype), which is rare in Asians, occurs in about 5% to 10% of Caucasians and in 2% of Negroids.

As another example, the N-acetyl transferase-1

35

(NAT1) and N-acetyl transferase-2 (NAT2) genes are located on chromosome 8q. Allelic variation in the NAT genes may contribute to variation in populations as to the susceptibility of individuals to development of breast
5 carcinoma, particularly in the context of exposure to compounds present in tobacco. NAT2 detoxifies or, conversely, activates aromatic amines found in tobacco smoke such as 4-aminobiphenyl. Both phenotypic assays and genotypic assays for NAT2 can be used to classify
10 individuals as rapid or slow acetylators. Genetic variants of the NAT2 gene have been cloned and 6 alleles at this locus have been identified: the F1 allele confers the fast acetylator phenotype. The distribution of NAT1 and NAT2 alleles differs widely between racial and ethnic groups.

As yet another example, the glutathione S-transferase-M1(GSTM1) gene is located on chromosome 1 and the gene for glutathione S-transferase-T1 (GSTT1) is located on chromosome 11q. A glutathione S-transferase-P1 (GSTP1) gene has also been identified.

15

20 Glutathione S-transferases detoxify a variety of carcinogens and cytotoxic drugs (for example, benzo(a)pyrene, monohalomethanes such as methyl chloride, ethylene oxide, pesticides, and solvents used in industry) by catalyzing the conjugation of a glutathione moiety to

the substrate. Allelic variation in the glutathione-Stransferase genes may contribute to variation in populations as to the susceptibility of individuals to development of breast carcinoma, particularly in the context of exposure to environmental toxins. Individuals

30 homozygous for deletions in the GSTM1, GSTT1, or GSTP1 genes may have a higher risk of cancer of the breast and other sites because of their impaired ability to metabolize and eliminate carcinogens.

GSTM1 is polymorphically expressed and 3 alleles at the GSTM1 locus have been identified: GSTM1-0

(homozygous deletion genotype), GSTM1a, and GSTM1b. The null allele (GSTM1-0) is present in about 38% to 67% of Caucasians and 22% to 35% of Negroids. GSTM is not expressed in breast tissue at high levels. Two

functionally different genotypes at the GSTT1 locus have been described: GSTT1-0 (homozygous deletion genotype) and GSTT1-1 (genotypes with 1 or 2 undeleted alleles). A polymorphism of the GSTP1 gene, A313G (changing codon 105 from Ile to Val), has been identified. The GSTT1-0 allele has been associated with accelerated age of first breast cancer diagnosis as compared with the GSTT1-1 allele.

Many other genes have been suggested to be involved in the development and/or progression of breast cancer, either as a result of gain of function or loss of function mutations, or as a result of normal allelic variation within different populations. A nonexhaustive list of such genes, each followed by the gene's chromosomal location, if known, follows: AMPH 7p14-p13; AMPHL (BIN1, SH3P9) 2q14; API4 (survivin, SVV) 17q25(?); ARHA (ARH12,

- 20 RhoA) 3p21.3; ARHC (RhoC) 1p21-p13; ATM (ATA, ATC) 11q22.3;
  BAG1 9p12; BARD1 2q34-q35; BCAR1 16q23.1; BCAR2; BCAR3
  (NSP2); BCAS1 (NABC1, AIBC1) 20q13.2-q13.3; BRCA1 17q21;
  BRCA2 13q12.3; CCND1 (D11S287E, Cyclin D, PRAD1) 11q13;
  CD44 (MDU3, HA, MDU2) 11pter-p13; CD9 (p24, MIC3, BA2)
- 25 12p13; CDKN1B (KIP1, P27) 12p13; CDKN2A (P16, INK4A, MTS1)
  9p21; COMT 22q11.2; COT (MAP3K8, TPL-2, EST) 10p11.2; CSK
  (c-src) 15q23-q25; CTSD (CPSD) 11p15.5; CYP17 10q24.3;
  CYP19 15q21.1; CYP1A1 (CYP1) 15q22-q24; CYP1B1 (GLC3A)
  2p22-p21; EFNB2 (EPLG5, LERK5, ephrin-B2) 13q33; EIF3S6
- 30 (INT6) 8q22-q23; EIF4E (EIF-4E) 4q21-q25; EMS1 11q13; ERBB2 (HER2, NEU) 17q11.2-q12; ERBB3 (HER3) 12q13; ESR1 (ESRA) 6q25.1; ESR2 (ESRB, ERBeta) 14q; FGF8 (AIGF) 10q24; GSTM1 (GST1, MU) 1p13.3; GSTP1 (FAEES3, GST3, PI) 11q13; GSTT1 22q11.23; HRAS 11p15.5; HSPB1 (HSP27) 7q;
- 35 HSPCA (HSP90A , HSPC1); HSPCB (HSP90B, HSPC2) 6p12; IGF1

12q22-q24.1; IGF1R (JTK13) 15q25-q26; IGF2 11p15.5; IL6 (IFNB2) 7p21; ING1 13q34; KISS1 (KiSS-1) 1q32; KLK3 (PSA, APS) 19q13; LASP1 (MLN50) 17q11-q21.3; LIBC 6q22; MAP2K4 (MKK4, SEK1, JNKK1) 17p11.2; MKI67 (Ki-67) 5 10q25-qter; MMP11 (STMY3, STR3) 22q11.2; MMP2 (CLG4A, CLG4, GELA) 16q13; MUC1 (PUM, PEM) 1q21; MYC (CMYC, C-MYC) 8q24.12-q24.13; NOTCH4 (INT3, NOTCH3) 6p21.3; PCNA 20p12; PI5 (maspin ) 18q21.3; PLAU (uPA , URK) 10q24; PSEN2 (D21S21, HPS2, BCEI) 21q22.3; RARB (NR1B2, HAP) 3p24; RB1 10 (Rb) 13q14.2; S100A4 (MTS1, P9KA, metastasin) 1q21; SLC22A1L (BWSCR1A, ORCTL2, IMPT1) 11p15.5; SNCG (BCSG1) 10q23.2-q23.3; SRD5A2 2p23; STIP1 (HSP70) 11?; STK11 (LKB1, PJS) 19p13.3; TFAP2A (AP2 , AP2TF) 6p24; TFAP2B (AP2B) 6p12; TFAP2C 20q13.2; TFF1 (D21S21, BCEI) 21q22.3; TGFBR2 3p22; TIMP2 17q25; TP53 (p53 , P53) 17q13.1; TPD52 (D52) 8q21; TPD52L1 (D53, hD53) 6q22-q23; TSG101 11p15.2-p15.1.

The etiology of non-cancerous disorders of the breast may also involve genetic factors. Such disorders include disorders of development, inflammatory diseases of the breast, fibrocystic changes, proliferative breast disease, and non-carcinoma tumors.

Disorders of development of the breast include

supernumerary nipples or breasts; accessory axillary breast
tissue; congenital inversion of the nipples; and
macromastia.

Inflammatory diseases of the breast include acute mastitis; periductal mastitis, also called recurrent subareolar absecess and sqamous metaplasia of the lactiferous ducts; mammary duct ectasia; fat necrosis; and granulomatous mastitis, including granulomatous lobular mastitis. Systemic granulomatous diseases that can affect the breast include Wegener granulomatosis and sarcoidosis.

Proliferative breast diseases include epithelial hyperplasia; sclerosing adenosis; and small duct
35 papillomas. Non-carcinoma tumors include stomal tumors

including fibroadenoma and phyllodes tumor, and sarcomas that include angiosarcoma, rhabdomyosarcoma, liposarcoma, leiomyosarcoma, chondrosarcoma and osteosarcoma. Other breast tumors include epithelial cell tumors including large duct papillomas.

The human genome-derived single exon nucleic acid probes and microarrays of the present invention are useful for predicting, diagnosing, grading, staging, monitoring and prognosing diseases of human breast, particularly those diseases with polygenic etiology. With each of the single exon probes described herein shown to be expressed at detectable levels in human breast cancer cells, and with about 2/3 of the probes identifying novel genes, the single exon microarrays of the present invention provide

15 exceptionally high informational content for such studies.

For example, diagnosis, grading, and/or staging of a disease can be based upon the quantitative relatedness of a patient gene expression profile to one or more reference expression profiles known to be characteristic of a given breast disease, or to specific grades or stages thereof.

In one embodiment, the patient gene expression profile is generated by hybridizing nucleic acids obtained directly or indirectly from transcripts expressed in the patient's breast to the genome-derived single exon microarray of the present invention. Reference profiles are obtained similarly by hybridizing nucleic acids from individuals with known disease. Methods for quantitatively relating gene expression profiles, without regard to the function of the protein encoded by the gene, are disclosed in WO 99/58720, incorporated herein by reference in its entirety.

In another approach, the genome-derived single exon probes and microarrays of the present invention can be used to interrogate genomic DNA, rather than pools of

expressed message; this latter approach permits predisposition to and/or prognosis of breast disease to be assessed through the massively parallel determination of altered copy number, deletion, or mutation in the patient's genome of exons known to be expressed in human breast. The algorithms set forth in WO 99/58720 can be applied to such genomic profiles without regard to the function of the protein encoded by the interrogated gene.

The utility is specific to the probe; at

sufficiently high hybridization stringency, which
stringencies are well known in the art — see Ausubel et al.
and Maniatis et al. — each probe reports the level of
expression of message specifically containing that ORF.

It should be appreciated, however, that the
probes of the present invention, for which expression in
the HBL 100 cells has been demonstrated are useful for both
measurement in the Breast and for survey of expression in
other tissues.

Significant among such advantages is the presence of probes for novel genes.

Examples 1 and 2, the methods described enable ORFs which are not present in existing expression databases to be identified. And the fewer the number of tissues in which the ORF can be shown to be expressed, the more likely the ORF will prove to be part of a novel gene: as further discussed in Example 2, ORFs whose expression was measurable in only a single of the tested tissues were represented in existing expression databases at a rate of only 11%, whereas 36% of ORFs whose expression was measurable in 9 tissues were present in existing expression databases, and fully 45% of those ORFs expressed in all ten tested tissues were present in existing expressed sequence databases.

Either as tools for measuring gene expression or

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tools for surveying gene expression, the genome-derived single exon probes of the present invention have significant advantages over the cDNA or EST-based probes that are currently available for achieving these utilities.

The genome-derived single exon probes of the present invention are useful in constructing genome-derived single exon microarrays; the genome-derived single exon microarrays, in turn, are useful devices for measuring and for surveying gene expression in the human.

Gene expression analysis using microarrays — conventionally using microarrays having probes derived from expressed message — is well-established as useful in the biological research arts (see Lockhart et al. Nature 405, 827-836).

Microarrays have been used to determine gene 15 expression profiles in cells in response to drug treatment (see, for example, Kaminski et al., "Global Analysis of Gene Expression in Pulmonary Fibrosis Reveals Distinct Programs Regulating Lung Inflammation and Fibrosis, " Proc. 20 Natl. Acad. Sci. USA 97(4):1778-83 (2000); Bartosiewicz et al., "Development of a Toxicological Gene Array and Quantitative Assessment of This Technology, " Arch. Biochem. Biophys. 376(1):66-73 (2000), viral infection (see for example, Geiss et al., "Large-scale Monitoring of Host Cell 25 Gene Expression During HIV-1 Infection Using cDNA Microarrays, " Virology 266(1):8-16 (2000)) and during cell processes such as differentiation, senescence and apoptosis (see, for example, Shelton et al., "Microarray Analysis of Replicative Senescence, " Curr. Biol. 9(17):939-45 (1999); 30 Voehringer et al., "Gene Microarray Identification of Redox and Mitochondrial Elements That Control Resistance or Sensitivity to Apoptosis, " Proc. Natl. Acad. Sci. USA 97(6):2680-5 (2000)).

Microarrays have also been used to determine 35 abnormal gene expression in diseased tissues (see, for

example, Alon et al., "Broad Patterns of Gene Expression Revealed by Clustering Analysis of Tumor and Normal Colon Tissues Probed by Oligonucleotide Arrays, " Proc. Natl. Acad. Sci. USA 96(12):6745-50 (1999); Perou et al., 5 "Distinctive Gene Expression Patterns in Human Mammary Epithelial Cells and Breast Cancers, Proc. Natl. Acad. Sci. USA 96(16):9212-7 (1999); Wang et al., "Identification of Genes Differentially Over-expressed in Lung Squamous Cell Carcinoma Using Combination of cDNA Subtraction and 10 Microarray Analysis, " Oncogene 19(12):1519-28 (2000); Whitney et al., "Analysis of Gene Expression in Multiple

Sclerosis Lesions Using cDNA Microarrays, " Ann. Neurol. 46(3):425-8 (1999)), in drug discovery screens (see, for example, Scherf et al., "A Gene Expression Database for the 15 Molecular Pharmacology of Cancer, " Nat. Genet. 24(3):236-44 (2000)) and in diagnosis to determine appropriate treatment strategies (see, for example, Sgroi et al., "In vivo Gene Expression Profile Analysis of Human Breast Cancer Progression, " Cancer Res. 59(22):5656-61 (1999)).

In microarray-based gene expression screens of pharmacological drug candidates upon cells, each probe provides specific useful data. In particular, it should be appreciated that even those probes that show no change in expression are as informative as those that do change, 25 serving, in essence, as negative controls.

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For example, where gene expression analysis is used to assess toxicity of chemical agents on cells, the failure of the agent to change a gene's expression level is evidence that the drug likely does not affect the pathway 30 of which the gene's expressed protein is a part. Analogously, where gene expression analysis is used to assess side effects of pharmacological agents - whether in lead compound discovery or in subsequent screening of lead compound derivatives - the inability of the agent to alter 35 a gene's expression level is evidence that the drug does

not affect the pathway of which the gene's expressed protein is a part.

WO 99/58720 provides methods for quantifying the relatedness of a first and second gene expression profile 5 and for ordering the relatedness of a plurality of gene expression profiles. The methods so described permit useful information to be extracted from a greater percentage of the individual gene expression measurements from a microarray than methods previously used in the art.

Other uses of microarrays are described in 10 Gerhold et al., Trends Biochem. Sci. 24(5):168-173 (1999) and Zweiger, Trends Biotechnol. 17(11):429-436 (1999); Schena et al.

The invention particularly provides genome-15 derived single-exon probes known to be expressed in HBL 100 The individual single exon probes can be provided in the form of substantially isolated and purified nucleic acid, typically, but not necessarily, in a quantity sufficient to perform a hybridization reaction.

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Such nucleic acid can be in any form directly hybridizable to the message that contains the probe's ORF, such as double stranded DNA, single-stranded DNA complementary to the message, single-stranded RNAcomplementary to the message, or chimeric DNA/RNA molecules 25 so hybridizable. The nucleic acid can alternatively or additionally include either nonnative nucleotides, alternative internucleotide linkages, or both, so long as complementary binding can be obtained. For example, probes can include phosphorothioates, methylphosphonates, 30 morpholino analogs, and peptide nucleic acids (PNA), as are described, for example, in U.S. Patent Nos. 5,142,047;

Usefully, however, such probes are provided in a form and quantity suitable for amplification, where the 35 amplified product is thereafter to be used in the

5,235,033; 5,166,315; 5,217,866; 5,184,444; 5,861,250.

hybridization reactions that probe gene expression.

Typically, such probes are provided in a form and quantity suitable for amplification by PCR or by other well known amplification technique. One such technique additional to PCR is rolling circle amplification, as is described, inter alia, in U.S. Patent Nos. 5,854,033 and 5,714,320 and international patent publications WO 97/19193 and WO 00/15779. As is well understood, where the probes are to be provided in a form suitable for amplification, the range of nucleic acid analogues and/or internucleotide linkages will be constrained by the requirements and nature of the amplification enzyme.

Where the probe is to be provided in form suitable for amplification, the quantity need not be sufficient for direct hybridization for gene expression analysis, and need be sufficient only to function as an amplification template, typically at least about 1, 10 or 100 pg or more.

packaged with amplification primers, either in a single composition that comprises probe template and primers, or in a kit that comprises such primers separately packaged therefrom. As earlier mentioned, the ORF-specific 5' primers used for genomic amplification can have a first common sequence added thereto, and the ORF-specific 3' primers used for genomic amplification can have a second, different, common sequence added thereto, thus permitting, in this embodiment, the use of a single set of 5' and 3' primers to amplify any one of the probes. The probe composition and/or kit can also include buffers, enzyme, etc., required to effect amplification.

As mentioned earlier, when intended for use on a genome-derived single exon microarray of the present invention, the genome-derived single exon probes of the present invention will typically average at least about

100, 200, 300, 400 or 500 bp in length, including (and typically, but not necessarily centered about) the ORF.
Furthermore, when intended for use on a genome-derived single exon microarray of the present invention, the genome-derived single exon probes of the present invention will typically not contain a detectable label.

When intended for use in solution phase
hybridization, however — that is, for use in a
hybridization reaction in which the probe is not first

bound to a support substrate (although the target may
indeed be so bound) — length constraints that are imposed
in microarray-based hybridization approaches will be
relaxed, and such probes will typically be labeled.

In such case, the only functional constraint that
dictates the minimum size of such probe is that each such
probe must be capable of specifically identifying in a
hybridization reaction the exon from which it is drawn. In
theory, a probe of as little as 17 nucleotides is capable
of uniquely identifying its cognate sequence in the human
genome. For hybridization to expressed message — a subset
of target sequence that is much reduced in complexity as
compared to genomic sequence — even fewer nucleotides are
required—for—specificity.

Therefore, the probes of the present invention

25 can include as few as 20, 25 or 50 bp or ORF, or more. In
particular embodiments, the ORF sequences are given in SEQ

ID NOS. 5,075 - 10,058, respectively, for probe SEQ ID NOS.

1 - 5,074. The minimum amount of ORF required to be
included in the probe of the present invention in order to

30 provide specific signal in either solution phase or
microarray-based hybridizations can readily be determined
for each of ORF SEQ ID NOS. 5,075 - 10,058 individually by
routine experimentation using standard high stringency
conditions.

Such high stringency conditions are described,

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inter alia, in Ausubel et al. and Maniatis et al. For microarray-based hybridization, standard high stringency conditions can usefully be 50% formamide, 5X SSC, 0.2 µg/µl poly(dA), 0.2 µg/µl human cot1 DNA, and 0.5 % SDS, in a humid oven at 42°C overnight, followed by successive washes of the microarray in 1X SSC, 0.2% SDS at 55°C for 5 minutes, and then 0.1X SSC, 0.2% SDS, at 55°C for 20 minutes. For solution phase hybridization, standard high stringency conditions can usefully be aqueous hybridization at 65°C in 6X SSC. Lower stringency conditions, suitable for cross-hybridization to mRNA encoding structurally- and functionally-related proteins, can usefully be the same as the high stringency conditions but with reduction in temperature for hybridization and washing to room temperature (approximately 25°C).

When intended for use in solution phase hybridization, the maximum size of the single exon probes of the present invention is dictated by the proximity of other expressed exons in genomic DNA: although each single exon probe can include intergenic and/or intronic material contiguous to the ORF in the human genome, each probe of the present invention will include portions of only one expressed exon.

Thus, each single exon probe will include no more
than about 25 kb of contiguous genomic sequence, more
typically no more than about 20 kb of contiguous genomic
sequence, more usually no more than about 15 kb, even more
usually no more than about 10 kb. Usually, probes that are
maximally about 5 kb will be used, more typically no more
than about 3 kb.

It will be appreciated that the Sequence Listing appended hereto presents, by convention, only that strand of the probe and ORF sequence that can be directly translated reading from 5' to 3' end. As would be well understood by one of skill in the art, single stranded

probes must be complementary in sequence to the ORF as present in an mRNA; it is well within the skill in the art to determine such complementary sequence. It will further be understood that double stranded probes can be used in both solution-phase hybridization and microarray-based hybridization if suitably denatured.

Thus, it is an aspect of the present invention to provide single-stranded nucleic acid probes that have sequence complementary to those described herein above and below, and double-stranded probes one strand of which has sequence complementary to the probes described herein.

The probes can, but need not, contain intergenic and/or intronic material that flanks the ORF, on one or both sides, in the same linear relationship to the ORF that the intergenic and/or intronic material bears to the ORF in genomic DNA. The probes do not, however, contain nucleic acid derived from more than one expressed ORF.

And when intended for use in solution
hybridization, the probes of the present invention can
usefully have detectable labels. Nucleic acid labels are
well known in the art, and include, inter alia, radioactive
labels, such as <sup>3</sup>H, <sup>32</sup>P, <sup>33</sup>P, <sup>35</sup>S, <sup>125</sup>I, <sup>131</sup>I; fluorescent
labels, such as Cy3, Cy5, Cy5, Cy7, SYBR®

Green and other labels described in Haugland,

Handbook of Fluorescent Probes and Research Chemicals, 7th
ed., Molecular Probes Inc., Eugene, OR (2000), or
fluorescence resonance energy transfer tandem conjugates
thereof; labels suitable for chemiluminescent and/or
enhanced chemiluminescent detection; labels suitable for

ESR and NMR detection; and labels that include one member
of a specific binding pair, such as biotin, digoxigenin, or
the like.

The probes, either in quantity sufficient for hybridization or sufficient for amplification, can be provided in individual vials or containers.

Alternatively, such probes can usefully be packaged as a plurality of such individual genome-derived single exon probes.

When provided as a collection of plural 5 individual probes, the probes are typically made available in amplifiable form in a spatially-addressable ordered set, typically one per well of a microtiter dish. Although a 96 well microtiter plate can be used, greater efficiency is obtained using higher density arrays.

If, as earlier mentioned, the ORF-specific 5' primers used for genomic amplification had a first common sequence added thereto, and the ORF-specific 3' primers used for genomic amplification had a second, different, common sequence added thereto, a single set of 15 5' and 3' primers can be used to amplify all of the probes from the amplifiable ordered set.

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Such collections of genome-derived single exon probes can usefully include a plurality of probes chosen for the common attribute of expression in the human HBL 100 cells.

In such defined subsets, typically at least 50, 60, 75, 80, 85, 90 or 95% or more of the probes will be chosen-by-their-expression-in-the-defined-tissue-or-celltype.

The single exon probes of the present invention, 25 as well as fragments of the single exon probes comprising selectively hybridizable portions of the probe ORF, can be used to obtain the full length cDNA that includes the ORF by (i) screening of cDNA libraries; (ii) rapid 30 amplification of cDNA ends ("RACE"); or (iii) other conventional means, as are described, inter alia, in Ausubel et al. and Maniatis et al.

It is another aspect of the present invention to provide genome-derived single exon nucleic acid microarrays 35 useful for gene expression analysis, where the term

"microarray" has the meaning given in the definitional section of this description, supra.

The invention particularly provides genomederived single-exon nucleic acid microarrays comprising a 5 plurality of probes known to be expressed in human HBL 100 cells. In preferred embodiments, the present invention provides human genome-derived single exon microarrays comprising a plurality of probes drawn from the group consisting of SEQ ID NOS.: 1 - 5,074.

When used for gene expression analysis, the 10 genome-derived single exon microarrays provide greater physical informational density than do the genome-derived single exon microarrays that have lower percentages of probes known to be expressed commonly in the tested tissue. 15 At a fixed probe density, for example, a given microarray surface area of the defined subset genome-derived single exon microarray can yield a greater number of expression measurements. Alternatively, at a given probe density, the same number of expression measurements can be obtained from 20 a smaller substrate surface area. Alternatively, at a fixed probe density and fixed surface area, probes can be provided redundantly, providing greater reliability in signal measurement for any given probe. Furthermore, with a higher percentage of probes known to be expressed in the 25 assayed tissue, the dynamic range of the detection means can be adjusted to reveal finer levels discrimination among the levels of expression.

Although particularly described with respect to their utility as probes of gene expression, particularly as probes to be included on a genome-derived single exon microarray, each of the nucleic acids having SEQ ID NOS.: 1 - 5,074 contains an open-reading frame, set forth respectively in SEQ ID NOS.: 5,075 - 10,058, that encodes a protein domain. Thus, each of SEQ ID NOS. 1 - 5,074 can be used, or that portion thereof in SEQ ID NOS. 5,075 - 10,058

used, to express a protein domain by standard in vitro recombinant techniques. See Ausubel et al. and Maniatis et al.

Additionally, kits are available commercially

that readily permit such nucleic acids to be expressed as protein in bacterial cells, insect cells, or mammalian cells, as desired (e.g., HAT Protein Expression & Purification System, ClonTech Laboratories, Palo Alto, CA; Adeno-X Expression System, ClonTech Laboratories, Palo Alto, CA; Protein Fusion & Purification (pMAL\*) System, New England Biolabs, Beverley, MA)

Furthermore, shorter peptides can be chemically synthesized using commercial peptide synthesizing equipment and well known techniques. Procedures are described, inter alia, in Chan et al. (eds.), Fmoc Solid Phase Peptide Synthesis: A Practical Approach (Practical Approach Series, (Paper)), Oxford Univ. Press (March 2000) (ISBN: 0199637245); Jones, Amino Acid and Peptide Synthesis (Oxford Chemistry Primers, No 7), Oxford Univ. Press (August 1992) (ISBN: 0198556683); and Bodanszky, Principles of Peptide Synthesis (Springer Laboratory), Springer Verlag (December 1993) (ISBN: 0387564314).

It is, therefore, another aspect of the invention to provide peptides comprising an amino acid sequence

25 translated from SEQ ID NOS.: 5,075 - 10,058. Such amino acid sequences are set out in SEQ ID NOS: 10,059 - 15,009. Any such recombinantly-expressed or synthesized peptide of at least 8, and preferably at least about 15, amino acids, can be conjugated to a carrier protein and used to generate antibody that recognizes the peptide. Thus, it is a further aspect of the invention to provide peptides that have at least 8, preferably at least 15, consecutive amino acids.

illustration and not by way of limitation.

### EXAMPLE 1

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Preparation of Single Exon Microarrays from ORFs Predicted 5 in Human Genomic Sequence

# Bioinformatics Results

All human BAC sequences in fewer than 10 pieces that had been accessioned in a five month period 10 immediately preceding this study were downloaded from This corresponds to ~2200 clones, totaling ~350 MB of sequence, or approximately 10% of the human genome.

After masking repetitive elements using the program CROSS\_MATCH, the sequence was analyzed for open 15 reading frames using three separate gene finding programs. The three programs predict genes using independent algorithmic methods developed on independent training sets: GRAIL uses a neural network, GENEFINDER uses a hidden Markoff model, and DICTION, a program proprietary to 20 Genetics Institute, operates according to a different heuristic. The results of all three programs were used to create a prediction matrix across the segment of genomic

The three gene finding programs yielded a range 25 of results. GRAIL identified the greatest percentage of genomic sequence as putative coding region, 2% of the data analyzed. GENEFINDER was second, calling 1%, and DICTION yielded the least putative coding region, with 0.8% of genomic sequence called as coding region.

The consensus data were as follows. GRAIL and GENEFINDER agreed on 0.7% of genomic sequence, GRAIL and DICTION agreed on 0.5% of genomic sequence, and the three programs together agreed on 0.25% of the data analyzed. That is, 0.25% of the genomic sequence was identified by 35 all three of the programs as containing putative coding

region.

ORFs predicted by any two of the three programs ("consensus ORFs") were assorted into "gene bins" using two criteria: (1) any 7 consecutive exons within a 25 kb window were placed together in a bin as likely contributing to a single gene, and (2) all ORFs within a 25 kb window were placed together in a bin as likely contributing to a single gene if fewer than 7 exons were found within the 25 kb window.

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#### PCR

The largest ORF from each gene bin that did not span repetitive sequence was then chosen for amplification, as were all consensus ORFs longer than 500 bp. This method approximated one exon per gene; however, a number of genes were found to be represented by multiple elements.

Previously, we had determined that DNA fragments fewer than 250 bp in length do not bind well to the aminomodified glass surface of the slides used as support substrate for construction of microarrays; therefore, amplicons were designed in the present experiments to approximate 500 bp in length.

gene bin, a 500 bp fragment of sequence centered on the ORF
25 was passed to the primer picking software, PRIMER3
(available online for use at
http://www-genome.wi.mit.edu/cgi-bin/primer/). A first
additional sequence was commonly added to each ORF-unique
5' primer, and a second, different, additional sequence was
30 commonly added to each ORF-unique 3' primer, to permit
subsequent reamplification of the amplicon using a single
set of "universal" 5' and 3' primers, thus immortalizing
the amplicon. The addition of universal priming sequences
also facilitates sequence verification, and can be used to
35 add a cloning site should some ORFs be found to warrant

further study.

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The ORFs were then PCR amplified from genomic DNA, verified on agarose gels, and sequenced using the universal primers to validate the identity of the amplicon 5 to be spotted in the microarray.

Primers were supplied by Operon Technologies (Alameda, CA). PCR amplification was performed by standard techniques using human genomic DNA (Clontech, Palo Alto, CA) as template. Each PCR product was verified by SYBR® 10 green (Molecular Probes, Inc., Eugene, OR) staining of agarose gels, with subsequent imaging by Fluorimager (Molecular Dynamics, Inc., Sunnyvale, CA). PCR amplification was classified as successful if a single band appeared.

The success rate for amplifying ORFs of interest directly from genomic DNA using PCR was approximately 75%. FIG. 5 graphs the distribution of predicted ORF (exon) length and distribution of amplified PCR products, with ORF length shown in red and PCR product length shown in blue 20 (which may appear black in the figure). Although the range of ORF sizes is readily seen to extend to beyond 900 bp, the mean predicted exon size was only 229 bp, with a median size of 150 bp (n=9498). With an average amplicon size of 475 ± 25 bp, approximately 50% of the average PCR 25 amplification product contained predicted coding region, with the remaining 50% of the amplicon containing either intron, intergenic sequence, or both.

Using a strategy predicated on amplifying about 500 bp, it was found that long exons had a higher PCR 30 failure rate. To address this, the bioinformatics process was adjusted to amplify 1000, 1500 or 2000 bp fragments from exons larger than 500 bp. This improved the rate of successful amplification of exons exceeding 500 bp, constituting about 9.2% of the exons predicted by the gene 35 finding algorithms.

Approximately 75% of the probes disposed on the array (90% of those that successfully PCR amplified) were sequence-verified by sequencing in both the forward and reverse direction using MegaBACE sequencer (Molecular Dynamics, Inc., Sunnyvale, CA), universal primers, and standard protocols.

Some genomic clones (BACs) yielded very poor PCR and sequencing results. The reasons for this are unclear, but may be related to the quality of early draft sequence or the inclusion of vector and host contamination in some submitted sequence data.

Although the intronic and intergenic material flanking coding regions could theoretically interfere with hybridization during microarray experiments, subsequent empirical results demonstrated that differential expression ratios were not significantly affected by the presence of noncoding sequence. The variation in exon size was similarly found not to affect differential expression ratios significantly; however, variation in exon size was observed to affect the absolute signal intensity (data not shown).

The 350 MB of genomic DNA was, by the above-described process, reduced to 9750 discrete probes, which were spotted in duplicate onto glass slides using commercially available instrumentation (MicroArray GenII Spotter and/or MicroArray GenIII Spotter, Molecular Dynamics, Inc., Sunnyvale, CA). Each slide additionally included either 16 or 32 E. coli genes, the average hybridization signal of which was used as a measure of background biological noise.

Each of the probe sequences was BLASTed against the human EST data set, the NR data set, and SwissProt GenBank (May 7, 1999 release 2.0.9).

One third of the probe sequences (as amplified)
35 produced an exact match (BLAST Expect ("E") values less

than 1 e<sup>-100</sup>) to either an EST (20% of sequences) or a known mRNA (13% of sequences). A further 22% of the probe sequences showed some homology to a known EST or mRNA (BLAST E values from 1 e<sup>-5</sup> to 1 e<sup>-99</sup>). The remaining 45% of the probe sequences showed no significant sequence homology to any expressed, or potentially expressed, sequences present in public databases.

All of the probe sequences (as amplified) were then analyzed for protein similarities with the SwissProt database using BLASTX, Gish et al., Nature Genet. 3:266 (1993). The predicted functional breakdowns of the 2/3 of probes identical or homologous to known sequences are presented in Table 1.

Table 1

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Function	of Predic	ted ORFs As	Deduced From Comparative
Sequence	Analysis		
Total	V6 chip	V7 chip	Function Predicted from Comparative Sequence Analysis
211	96	115	Receptor
120	43	77	Zinc Finger
30	11	19	Homeobox
25	9	16	Transcription Factor
17	11	7	Transcription
118	57	61	Structural
95	39	56	Kinase
36	18	18	Phosphatase
83	31	52	Ribosomal
45	19	26	Transport
21	17	14	Growth Factor
17	12	5	Cytochrome
50	33	17	Channel

As can be seen, the two most common types of genes were transcription factors and receptors, making up 2.2% and 1.8% of the arrayed elements, respectively.

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### EXAMPLE 2

Gene Expression Measurements From Genome-Derived Single Exon Microarrays

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The two genome-derived single exon microarrays prepared according to Example 1 were hybridized in a series of simultaneous two-color fluorescence experiments to (1)

15 Cy3-labeled cDNA synthesized from message drawn individually from each of brain, heart, liver, fetal liver, placenta, lung, bone marrow, HeLa, BT 474, or HBL 100 cells cells, and (2) Cy5-labeled cDNA prepared from message pooled from all ten tissues and cell types, as a control in each of the measurements. Hybridization and scanning were carried out using standard protocols and Molecular Dynamics equipment.

Briefly, mRNA samples were bought from commercial sources (Clontech, Palo Alto, CA and Amersham Pharmacia

25 Biotech (APB)). Cy3-dCTP and Cy5-dCTP (both from APB) were incorporated during separate reverse transcriptions of 1 µg of polyA<sup>+</sup> mRNA performed using 1 µg oligo(dT)12-18 primer and 2 µg random 9mer primers as follows. After heating to 70°C, the RNA:primer mixture was snap cooled on ice. After snap cooling on ice, added to the RNA to the stated final concentration was: 1X Superscript II buffer, 0.01 M DTT, 100µM dATP, 100 µM dGTP, 100 µM dTTP, 50 µM dCTP, 50 µM Cy3-dCTP or Cy5-dCTP 50 µM, and 200 U Superscript II enzyme. The reaction was incubated for 2 hours at 42°C.

35 After 2 hours, the first strand cDNA was isolated by adding

1 U Ribonuclease H, and incubating for 30 minutes at 37°C. The reaction was then purified using a Qiagen PCR cleanup column, increasing the number of ethanol washes to 5. Probe was eluted using 10 mM Tris pH 8.5.

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Using a spectrophotometer, probes were measured for dye incorporation. Volumes of both Cy3 and Cy5 cDNA corresponding to 50 pmoles of each dye were then dried in a Speedvac, resuspended in 30 µl hybridization solution containing 50% formamide, 5% SSC, 0.2 µg/µl poly(dA), 0.2 10 µg/µl human cot1 DNA, and 0.5 % SDS.

Hybridizations were carried out under a coverslip, with the array placed in a humid oven at 42°C overnight. Before scanning, slides were washed in 1X SSC, 0.2% SDS at 55°C for 5 minutes, followed by 0.1% SSC, 0.2% 15 SDS, at 55°C for 20 minutes. Slides were briefly dipped in water and dried thoroughly under a gentle stream of nitrogen.

Slides were scanned using a Molecular Dynamics Gen3 scanner, as described. Schena (ed.), Microarray 20 Biochip: Tools and Technology, Eaton Publishing Company/BioTechniques Books Division (2000) (ISBN: 1881299376)....

Although the use of pooled cDNA as a reference permitted the survey of a large number of tissues, it 25 attenuates the measurement of relative gene expression, since every highly expressed gene in the tissue/cell typespecific fluorescence channel will be present to a level of at least 10% in the control channel. Because of this fact, both signal and expression ratios (the latter hereinafter, 30 "expression" or "relative expression") for each probe were normalized using the average ratio or average signal, respectively, as measured across the whole slide.

Data were accepted for further analysis only when signal was at least three times greater than biological 35 noise, the latter defined by the average signal produced by

the E. coli control genes.

The relative expression signal for these probes was then plotted as function of tissue or cell type, and is presented in FIG. 6.

FIG. 6 shows the distribution of expression across a panel of ten tissues. The graph shows the number of sequence-verified products that were either not expressed ("0"), expressed in one or more but not all tested tissues ("1" - "9"), and expressed in all tissues tested ("10").

Of 9999 arrayed elements on the two microarrays (including positive and negative controls and "failed" products), 2353 (51%) were expressed in at least one tissue or cell type. Of the gene elements showing significant signal — where expression was scored as "significant" if the normalized Cy3 signal was greater than 1, representing signal 5-fold over biological noise (0.2) — 39% (991) were expressed in all 10 tissues. The next most common class (15%) consisted of gene elements expressed in only a single tissue.

The genes expressed in a single tissue were further analyzed, and the results of the analyses are compiled in FIG. 7.

FIG. 7A is a matrix presenting the expression of
all verified sequences that showed expression greater than
3 in at least one tissue. Each clone is represented by a
column in the matrix. Each of the 10 tissues assayed is
represented by a separate row in the matrix, and relative
expression of a clone in that tissue is indicated at the
respective node by intensity of green shading, with the
intensity legend shown in panel B. The top row of the
matrix ("EST Hit") contains "bioinformatic" rather than
"physical" expression data — that is, presents the results
returned by query of EST, NR and SwissProt databases using
the probe sequence. The legend for "bioinformatic

expression" (i.e., degree of homology returned) is presented in panel C. Briefly, white is known, black is novel, with gray depicting nonidentical with significant homology (white: E values < 1e-100; gray: E values from 1e-5 05 to 1e-99; black: E values > 1e-05).

As FIG. 7 readily shows, heart and brain were demonstrated to have the greatest numbers of genes that were shown to be uniquely expressed in the respective tissue. In brain, 200 uniquely expressed genes were identified; in heart, 150. The remaining tissues gave the following figures for uniquely expressed genes: liver, 100; lung, 70; fetal liver, 150; bone marrow, 75; placenta, 100; HeLa, 50; HBL, 100; and BT474, 50.

"novel" genes among those that were up-regulated in only one tissue, as compared with those that were down-regulated in only one tissue. In fact, it was found that ORFs whose expression was measurable in only a single of the tested tissues were represented in sequencing databases at a rate of only 11%, whereas 36% of the ORFs whose expression was measurable in 9 of the tissues were present in public databases. As for those ORFs expressed in all ten tissues, fully 45% were present in existing expressed sequence databases. These results are not unexpected, since genes expressed in a greater number of tissues have a higher likelihood of being, and thus of having been, discovered by EST approaches.

# Comparison of Signal from Known and Unknown Genes

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The normalized signal of the genes found to have high homology to genes present in the GenBank human EST database were compared to the normalized signal of those genes not found in the GenBank human EST database. The data are shown in FIG. 8.

FIG. 8 shows the normalized Cy3 signal intensity

for all sequence-verified products with a BLAST Expect
("E") value of greater than 1e-30 (designated "unknown")
upon query of existing EST, NR and SwissProt databases, and
shows in blue the normalized Cy3 signal intensity for all
sequence-verified products with a BLAST Expect value of
less than 1e-30 ("known"). Note that biological background
noise has an averaged normalized Cy3 signal intensity of
0.2.

As expected, the most highly expressed of the
10 ORFs were "known" genes. This is not surprising, since
very high signal intensity correlates with very commonlyexpressed genes, which have a higher likelihood of being
found by EST sequence.

However, a significant point is that a large

15 number of even the high expressers were "unknown". Since
the genomic approach used to identify genes and to confirm
their expression does not bias exons toward either the 3'
or 5' end of a gene, many of these high expression genes
will not have been detected in an end-sequenced cDNA

20 library.

The significant point is that presence of the

gene\_in\_an EST database\_is\_not a prerequisite for
incorporation into a genome-derived microarray, and
further, that arraying such "unknown" exons can help to
assign function to as-yet undiscovered genes.

# Verification of Gene Expression

To ascertain the validity of the approach described above to identify genes from raw genomic

sequence, expression of two of the probes was assayed using reverse transcriptase polymerase chain reaction (RT PCR) and northern blot analysis.

Two microarray probes were selected on the basis of exon size, prior sequencing success, and tissue-specific gene expression patterns as measured by the microarray

experiments. The primers originally used to amplify the two respective ORFs from genomic DNA were used in RT PCR against a panel of tissue-specific cDNAs (Rapid-Scan gene expression panel 24 human cDNAs) (OriGene Technologies, Inc., Rockville, MD).

Sequence AL079300\_1 was shown by microarray hybridization to be present in cardiac tissue, and sequence AL031734\_1 was shown by microarray experiment to be present in placental tissue (data not shown). RT-PCR on these two sequences confirmed the tissue-specific gene expression as measured by microarrays, as ascertained by the presence of a correctly sized PCR product from the respective tissue type cDNAs.

Clearly, all microarray results cannot, and
indeed should not, be confirmed by independent assay
methods, or the high throughput, highly parallel advantages
of microarray hybridization assays will be lost. However,
in addition to the two RT-PCR results presented above, the
observation that 1/3 of the arrayed genes exist in
expression databases provides powerful confirmation of the
power of our methodology — which combines bioinformatic
prediction with expression confirmation using genomederived single exon microarrays — to identify novel genes
from raw genomic data.

To verify that the approach further provides correct characterization of the expression patterns of the identified genes, a detailed analysis was performed of the microarrayed sequences that showed high signal in brain.

25

For this latter analysis, sequences that showed

high (normalized) signal in brain, but which showed very
low (normalized) signal (less than 0.5, determined to be
biological noise) in all other tissues, were further
studied. There were 82 sequences that fit these criteria,
approximately 2% of the arrayed elements. The 10 sequences
showing the highest signal in brain in microarray

hybridizations are detailed in Table 2, along with assigned function, if known or reasonably predicted.

Table 2

ble 2		6 11 2 200	+ Wighly	
		of the Mos		,
Expressed G	enes Exp	ressed On	y III Diai	•
Microarray Sequence Name	Normal ized Signal	Expressi on Ratio	to EST present in	as described by
· .			GenBank	
AP000217-1	5.2	+7.7	High	S-100 protein, b-chain, Ca <sup>2+</sup> binding protein expressed in central nervous system
AP000047-1	2.3		High	Unknown Function
AC006548-9	1.7		High	Similar to mouse_membrane
				glyco-protein M6, expressed in central nervous system
AC007245-5	1.5		High	Similar to amphiphysin, a synaptic vesicle- associated protein. Ref 21
L44140-4	1.2	+2.0	High	Endothelial actin-binding

	Т			protein found
1				in nonmuscle
				filamin
		+3.5	High	Protein
AC004689-9	1.2 .	+3.5	1119	Phosphatase
			,	PP2A, neuronal/
				downregulates
				activated
·				protein kinases
AL031657-1	1.2	+3.0	High	Unknown
ALUSTOSTAL	1.2			function/
		,		Contains the
				anhyrin motif,
				a common
				protein
				sequence motif
AC009266-2	1.1	+3.7	Low	Low homology to
				the
	•		,	Synaptotagmin I
		,		protein in
				rat/present at
				low levels
				throughout rat
				brain
AP000086-1	1.0	+2.7	Low	Unknown, very
				poor homology
		<b>!</b> ,		to collagen
AC004689-3	1.0		High	Protein
		}		Phosphatase
				PP2A, neuronal/
•	·			downregulates
			,	activated
				protein kinases

Of the ten sequences studied by these latter

confirmatory approaches, eight were previously known. Of these eight, six had previously been reported to be important in the central nervous system or brain. The exon giving the highest signal (AP00217-1) was found to be the gene encoding an S100B Ca<sup>2+</sup> binding protein, reported in the literature to be highly and uniquely expressed in the central nervous system. Heizmann, Neurochem. Res. 9:1097 (1997).

A number of the brain-specific probe sequences

(including AC006548-9, AC009266-2) did not have homology to any known human cDNAs in GenBank but did show homology to rat and mouse cDNAs. Sequences AC004689-9 and AC004689-3 were both found to be phosphatases present in neurons (Millward et al., Trends Biochem. Sci. 24(5):186-191

(1999)). Two microarray sequences, AP000047-1 and AP000086-1 have unknown function, with AP000086-1 being absent from GenBank. Functionality can now be narrowed down to a role in the central nervous system for both of these genes, showing the power of designing microarrays in this fashion.

Next, the function of the chip sequences with the
highest (normalized) signal intensity in brain, regardless
of expression in other tissues, was assessed. In this
latter analysis, we found expression of many more common

25 genes, since the sequences were not limited to those
expressed only in brain. For example, looking at the 20
highest signal intensity spots in brain, 4 were similar to
tubulin (AC00807905; AF146191-2; AC007664-4; AF14191-2), 2
were similar to actin (AL035701-2; AL034402-1), and 6 were

30 found to be homologous to glyceraldehyde-3-phosphate
dehydrogenase (GAPDH) (AL035604-1; Z86090-1; AC006064-L,
AC006064-K; AC035604-3; AC006064-L). These genes are often
used as controls or housekeeping genes in microarray
experiments of all types.

Other interesting genes highly expressed in brain

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were a ferritin heavy chain protein, which is reported in the literature to be found in brain and liver (Joshi et al., J. Neurol. Sci. 134(Suppl):52-56 (1995)), a result duplicated with the array. Other highly expressed chip sequences included a translation elongation factor 10 (AC007564-4), a DEAD-box homolog (AL023804-4), and a Y-chromosome RNA-binding motif (Chai et al., Genomics 49(2):283-89 (1998)) (AC007320-3). A low homology analog (AP00123-1/2) to a gene, DSCR1, thought to be involved in trisomy 21 (Down's syndrome), showed high expression in both brain and heart, in agreement with the literature (Fuentes et al., Mol. Genet. 4(10):1935-44 (1995)).

As a further validation of the approach, we selected the BAC AC006064 to be included on the array.

This BAC was known to contain the GAPDH gene, and thus could be used as a control for the ORF selection process. The gene finding and exon selection algorithms resulted in choosing 25 exons from BAC AC006064 for spotting onto the array, of which four were drawn from the GAPDH gene. Table 3 shows the comparison of the average expression ratio for the 4 exons from BAC006064 compared with the average expression ratio for 5 different dilutions of a commercially available GAPDH cDNA (Clontech).

#### 25 Table 3

Compar	ison of Expression R	atio, for each
tissue, of GAPDH		
-	AC006064 (n = 4)	Control (n = 5)
Bone Marrow	-1.81 ± 0.11	-1.85 ± 0.08
Brain	-1.41 ± 0.11	-1.17 ± 0.05
BT474	1.85 ± 0.09	1.66 ± 0.12
Fetal Liver	-1.62 ± 0.07	-1.41 ± 0.05
HBL100	1.32 ± 0.05	2.64 ± 0.12

Heart	1.16 ± 0.09	1.56 ± 0.10
	1.11 ±0.06	1.30 ± 0.15
НеБа	-1.62 ± 0.22	-2.07 ±
Liver		$-3.75 \pm 0.21$
Lung	-4.95 ± 0.93	$-3.52 \pm 0.43$
Placenta	-3.56 ± 0.25	-3.32 1 0.43

Each tissue shows excellent agreement between the experimentally chosen exons and the control, again demonstrating the validity of the present exon mining approach. In addition, the data also show the variability of expression of GAPDH within tissues, calling into question its classification as a housekeeping gene and utility as a housekeeping control in microarray experiments.

### EXAMPLE 3

Representation of Sequence and Expression Data as a "Mondrian"

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as above-described, a plethora of information was accumulated, including full clone sequence, probe sequence within the clone, results of each of the three gene finding programs, EST information associated with the probe sequences, and microarray signal and expression for multiple tissues, challenging our ability to display the information.

Accordingly, we devised a new tool for visual display of the sequence with its attendant annotation which, in deference to its visual similarity to the paintings of Piet Mondrian, is hereinafter termed a "Mondrian". FIGS. 3 and 4 present the key to the information presented on a Mondrian.

FIG. 9 presents a Mondrian of BAC AC008172 (bases

25,000 to 130,000 shown), containing the carbamyl phosphate synthetase gene (AF154830.1). Purple background within the region shown as field 81 in FIG. 3 indicates all 37 known exons for this gene.

As can be seen, GRAIL II successfully identified 27 of the known exons (73%), GENEFINDER successfully identified 37 of the known exons (100%), while DICTION identified 7 of the known exons (19%).

Seven of the predicted exons were selected for physical assay, of which 5 successfully amplified by PCR and were sequenced. These five exons were all found to be from the same gene, the carbamyl phosphate synthetase gene (AF154830.1).

The five exons were arrayed, and gene expression

15 measured across 10 tissues. As is readily seen in the

Mondrian, the five chip sequences on the array show

identical expression patterns, elegantly demonstrating the
reproducibility of the system.

FIG. 10 is a Mondrian of BAC AL049839. We selected 12 exons from this BAC, of which 10 successfully sequenced, which were found to form between 5 and 6 genes. Interestingly, 4 of the genes on this BAC are protease inhibitors. Again, these data elegantly show that exons selected from the same gene show the same expression 25 patterns, depicted below the red line. From this figure, it is clear that our ability to find known genes is very good. A novel gene is also found from 86.6 kb to 88.6 kb, upon which all the exon finding programs agree. We are confident we have two exons from a single gene since they 30 show the same expression patterns and the exons are proximal to each other. Backgrounds in the following colors indicate a known gene (top to bottom): red = kallistatin protease inhibitor (P29622); purple = plasma serine protease inhibitor (P05154); 35 turquoise =  $\alpha$ 1 anti-chymotrypsin (P01011); mauve = 40S

ribosomal protein (P08865). Note that chip sequence 8 and 12 did not sequence verify.

### 5 EXAMPLE 4

Genome-Derived Single Exon Probes Useful For Measuring Human Gene Expression

The protocols set forth in Examples 1 and 2,

10 supra, were applied to additional human genomic sequence as

it became newly available in GenBank to identify unique

exons in the human genome that could be shown to be

expressed at significant levels in HBL 100 cells.

These unique exons are within longer probe

sequences. Each probe was completely sequenced on both strands prior to its use on a genome-derived single exon microarray; sequencing confirms the exact chemical structure of each probe. An added benefit of sequencing is that it placed us in possession of a set of single base-incremented fragments of the sequenced nucleic acid, starting from the sequencing primer 3' OH. (Since the single exon probes were first obtained by PCR amplification from genomic DNA, we were of course additionally in possession of an even larger set of single base incremented fragments of each of the 5,074 single exon probes, each fragment corresponding to an extension product from one of the two amplification primers.)

The structures of the 5,074 unique single exon probes are clearly presented in the Sequence Listing as SEQ 30 ID Nos.: 1 - 5,074. The 16 nt 5' primer sequence and 16 nt 3' primer sequence present on the amplicon are not included in the sequence listing. The sequences of the exons present within each of these probes is presented in the Sequence Listing as SEQ ID Nos.: 5,075 - 10,058, respectively. It will be noted that some amplicons have

more than one exon, some exons are contained in more than one amplicon.

As detailed in Example 2, expression was
demonstrated by disposing the amplicons as single exon
probes on nucleic acid microarrays and then performing twocolor fluorescent hybridization analysis; significant
expression is based on a statistical confidence that the
signal is significantly greater than negative biological
control spots. The negative biological control is formed
from spotted DNA sequences from a different species. Here,
32 sequences from E.Coli were spotted in duplicate to give
a total of 64 spots.

For each hybridisation (each slide, each colour) the median value of the signal from all of the spots is determined. The normalised signal value is the arithmetic mean of the signal from duplicate spots divided by the population median.

Control spots are eliminated if there is more that a five-fold difference between each one of the duplicate spots raw signals.

The median of the signal from the remaining control spots is calculated and all subsequent calculations are done with normalised signals.

Control spots having a signal of greater than

25 median + 2.4 (the value 2.4 is roughly 12 times the
observed standard deviation of control spot populations)
are eliminated. Spots with such high signals are considered
to be "outliers".

The mean and standard deviation of the modified control spot populations are calculated.

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The mean + 3x the standard deviation (mean + (3\*SD)) is used as the signal threshold qualifier for that particular hybridisation. Thus, individual thresholds are determined for each channel and each hybridisation.

This means that, assuming that the data is

distributed normally, there is a 99% confidence that any signal exceeding the threshold is significant.

The probes and their expression data are presented in Table 4, set forth respectively in Example 5. 5 Example 5 presents the subset of probes that is significantly expressed in the human HBL 100 cells and thus presents the subset of probes that was recognized to be useful for measuring expression of their cognate genes in human HBL 100 cells.

The sequence of each of the exon probes identified by SEQ ID NOS.: 5,075 - 10,058 was individually used as a BLAST (or, for SWISSPROT, BLASTX) query to identify the most similar sequence in each of dbEST, SwissProt (BLASTX), and NR divisions of GenBank. Because 15 the query sequences are themselves derived from genomic sequence in GenBank, only nongenomic hits from NR were scored.

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The smallest in value of the BLAST (or BLASTX) expect ("E") scores for each query sequence across the 20 three database divisions was used as a measure of the "expression novelty" of the probe's ORF. Table 4 is sorted in descending order based on this measure, reported as "Most Similar (top) Hit BLAST E Value". Those sequences for which no "Hit E Value" is listed are those exons which were 25 found to have no similar sequences.

As sorted, Table 4 thus lists its respective probes (by "AMPLICON SEQ ID NO .: " and additionally by the SEQ ID NO:. of the exon contained within the probe: "EXON SEQ ID NO.: ") from least similar to sequences known to be 30 expressed (i.e., highest BLAST E value), at the beginning of the table, to most similar to sequences known to be expressed (i.e., lowest BLAST E value), at the bottom of the table.

Table 4 further provides, for each listed probe, 35 the accession number of the database sequence that yielded

the "Most Similar (top) Hit BLAST E Value", along with the name of the database in which the database sequence is found ("Top Hit Database Source").

Table 4 further provides SEQ ID NOS.

5 corresponding to the predicted amino acid sequences where they have been determined for the probe and exon nucleotide sequences. These are set out as PEPTIDE SEQ ID NOS .: . The peptide sequences for a given exon are predicted as follows: Since each chip exon is a consensus sequence drawn 10 from predictions from various exon finding programs (i.e. Grail, GeneFinder and GenScan), the multiple initial ORFs are first determined in a uniform way according to each prediction. In particular, the reading frame for predicting the first amino acid in the peptide sequence always starts 15 with the first base of any codon and ends with the last base of non-termination codon. Next, for each strand of the exon, initial ORFs are merged into one or more final ORFs in an exhaustive process based on the following criteria: 1) the merging ORFs must be overlapping, and 2) the merging 20 ORFs must be in the same frame.

The Sequence Listing, which is a superset of all

of the data presented in Table 4, further includes, for
each probe, the most similar hit, with accession number and
BLAST E value, from the each of the three queried

25 databases.

Table 4 further lists, for each probe, a portion of the descriptor for the top hit ("Top Hit Descriptor") as provided in the sequence database. For those ORFs that are similar in sequence, but nonidentical to known sequences (e.g., those with BLAST E values between about 1e-05 and 1e-100), the descriptor reveals the likely function of the protein encoded by the probe's ORF.

Using BLAST E value cutoffs of 1e-05 (i.e., 1 x  $10^{-5}$ ) and 1e-100 (i.e., 1 x  $10^{-100}$ ) as evidence of similarity to sequences known to be expressed is of course arbitrary:

in Example 2, supra, a BLAST E value of 1e-30 was used as the boundary when only two classes were to be defined for analysis (unknown, >1e-30; known <1e-30) (see also FIG. 8). Furthermore, even when the "Most Similar (Top) Hit BLAST E Value" is low, e.g., less than about 1e-100 — which is probative evidence that the query sequence has previously been shown to be expressed — the top hit is highly unlikely exactly to match the probe sequence.

First, such expression entries typically will not have the intronic and/or intergenic sequence present within the single exon probes listed in the Table. Second, even the ORF itself is unlikely in such cases to be present identically in the databases, since most of the EST and mRNA clones in existing databases include multiple exons, without any indication of the location of exon boundaries.

As noted, the data presented in Table 4 represent a proper subset of the data present within the attached sequence listing. For each amplicon probe (SEQ ID NOs.: 1 - 5,074) and probe exon (SEQ ID NOs.: 5,075 - 10,058, respectively), the sequence listing further provides, through iterated annotation fields <220> and <223>:

- (a) the accession number of the BAC from which the sequence was derived ("MAP TO"), thus providing a link to the chromosomal map location and other information about the genomic milieu of the probe sequence;
  - (b) the most similar sequence provided by BLAST query of the EST database, with accession number and BLAST E value for the "hit";
- (c) the most similar sequence provided by BLAST query of the GenBank NR database, with accession number and BLAST E value for the "hit"; and
  - (d) the most similar sequence provided by BLASTX query of the SWISSPROT database, with accession number and BLAST E value for the "hit".

## EXAMPLE 5

Genome-Derived Single Exon Probes Useful For Measuring
Expression of Genes in Human HBL 100 cells

Table 4 (209 pages) presents expression, homology, and functional information for the genome-derived single exon probes that are expressed significantly in human HBL 100 cells, a hormone sensitive human breast cancer cell line.

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Page 1 of 209 Table 4

		_	-	-	_	_	~	_	┰	_	_	_	_	_	_	_	7	_	-	_	P	Œ	7	-	<del>(</del> 1	H	<del></del>				H	Н	- Fa
Single Exon Probes Expressed in HBL100 Celis	Top Hit Descriptor																										Dengue virus type 3 membrane protein (prMM)/envelope glycoprotein (E) polyprotein mRNA, partial cds	Dengue virus Ivoe 3 membrane protein (pr/M/M/verwelope alvocondein (E) polvoraten mRNA pertial cds	Mus musculus AT3 gene for antithrombin, complete cds	Homo sepiens Insulin receptor substrate 1 (IRS1) mRNA	Lycopersicon esculentum Mill. GTPase (SAR2) mRNA, complete cds	Lycopersicon esculentum Mill. GTPase (SAR2) mRNA, complete cds	Homo sapiens DESC1 protein (DESC1), mRNA
con Probes E	Top Hit Database Source				_										_		_									·	_ <del>_</del>	<u>_</u>		11	NT		
Single E	Top Hit Acession No.																										L11433.1	11433.1	5.1	5031804 NT		L12051.1	7661557 NT
}_    - 	Most Similar (Top) Hit BLAST E Value																										9.4E+00	9.4E+00		_	7.2E+00	7.2E+00	5.8E+00
	Expression Signel	4.37	8.5	2.62	. 8.84	3.1	4.71	2.18	1.16	7.06	1.12	1.75	1.88	4.79	3.32	1.09	12.48	1.38	1.16	1.1	1.73	6.13	1.28	1.1	4.14	1.25	0.93	0.83	3.02	2.23	3.48	3.48	0.71
	ORF SEQ ID NO:	10501	10937		11324	11636	11653	11758	11781	11789	11928	12018	12197	12315	13148	13394	13471		13586		14058	14117		14657	14829	14837	12672	12673	12888	10491	12946	12947	
	SEQ ID NO:	5484	9689	6038	6282	6574	6592	6683	6705	6712	6839	6918	7082	7193	8128	8374	8445	8489	8580	8846	8906	9134	9204	9674	9858	9867	7558	7658	7873	5475			8453
	Probe SEQ ID NO:	447	878	1028	1283	1577	1596	1687	1710	1717	1850	1933	2101	2216	3112	3366	3437	3481	3573	3844	4074	4139	4211	4689	4879	4888	2692	2595	2853	437	2909	2909	3445

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Table 4
Single Exon Probes Expressed in HBL100 Cells

						_		_				_	_		_		_	7	-			Ť	"		7		4	7	4	4	٠.	۳ t	
Top Hit Descriptor	Homo sepiens fatty acid omega-hydroxylase CYP4A11 (CYP4A11) gene, complete cds	UI-H-BI3-84-9-08-0-UI:s1 NCI_CGAP_Sub5 Homo sapiens CUNA Glorie IMAGE:2734350 3	Loren emisone n27D-kidel (DOKDEL) mRNA, complete cds	Users and an expended (DOKDEL) mRNA, complete cds	Tromo seprens prze owac (2000-20) m	Ulycoagus cuinciles ner in the control of the contr	PUTATIVE RRINA METHYLI KANSFERASE STOT	R.norvegicus miKNA for collegen alpha i type i	R.norvegicus mRNA for collegen alpha i type i	INTSCBAT NOT CGAP_COT NOTICE SEPARATE DEHYDROGENASE, LIVER (HUMAN): GLYCERALDEHYDE 3-PHOSPHATE DEHYDROGENASE, LIVER (HUMAN):	hisage, xi NCI_CGAP_GU1 Hamo sapiens cuna dana invariation of summing agriculture allocations and summing and summing of yeeral Delydon and summing and summing and year summing and summi	PROTEIN B8 PRECURSOR	Syne-procedus sp. PCC7942 copper transporting P-ATPase (ctaA) and ATP synthese epsilon suburit	(apE) genes, complete cds	Synechococcus sp. PCC7942 copper transporting P-ATPass (clas) and ATP synitiate epsium euroning	(alpE) genes, complete cds	LEVANSUCRASE (BETA-D-FRUCTOFURANOSYL TRANSFERASE) (SUCRUSE OF RUCK LOSTE	TRANSFERASE)	Homo sapiens chightosame 21 segment 102.000	0243/NUS.X1 Soares Arithalting John Complete Cds	Indian septemble of the complete comple	Museum STRGalNAcili gene axon 2	The Hardward of and	2.725/11 1.1 Spares (she heart NDHH19W Home septens cDNA clone IMAGE:341689 5' similar to	gb:D29805 N-ACETYLACTOSAMINE SYNTHASE (HUMAN);	60218609571 NIH_MGC_45 Homo saplens cDNA clone IMAGE:4310591 3	Hamo saplens proliferation-associated SNF2-like protein (SNARCA) mixty, ampliate the	Homo sapiens proliferation associated SNF2-like protein (SMARCAB) miXNA, complete our	Mus musculus ST6GalNAcili gene, exon 2	Mus musculus ST6GalNAcill gene, exon 2	Rethis noveoleus fun dimerization protein 2 (jdp-2) mRNA, complete cds	Chlamydophila pneumonlae AR39, section 32 of 94 of the complete genome	
Top Hit Database Source		ECT HIMAN	Т		F	_	SWISSPROT	T	NT	EST_HUMAN	EST HUMAN	SWISSPROT		¥		_ <b>F</b>		SWISSPROT	NT	EST HUMAN	Z	- !	- !	Ž	EST HUMAN	EST HUMAN	LN	Ę	17		Į.	2	
<u>§</u>	본	ľ		Ž	Ż	Ż	S	N	Z	Ü		S	+	_ <u>z</u>	t		T	ő		٦				Ī							T	٦	7
Top Hit Acession No.	F208532.1	14/4/00/00 4	2.1E+00 AW 448500.1	2.0E+00 AF180527.1	\F180527.1	4F204927.1	>25582	278279.1	2.0E+00 Z78279.1	2.0E+00 AW664496.1	2 OF +00 AW664486 1	P21004		1 BF+00 004356.1		1104358.1	יבי	Q60114	-	1	~ 1		_	X98373.1	W 58428 1						711344.1	0 053449.1	JAE002201.2
Most Similar (Top) Hit BLAST E Value	2.1E+00 AF	100:11:0	2.1=+00/	2.0E+00 /	2.0E+00 /	2.0E+00	2.0E+00	2.0E+00	2.0E+00	2.0E+00	205+00	1 BE +00 P21004	3	1 8F+00		4 RF+00	20:1	1.7E+00	1.7E+00	1.7E+00	1.6E+00		1.6E+00	1.6E+00	0100								1.5E+00
Expression Signal	12		0.83	2.45	2.45	1.07	3.99	8.21	8.21	1.95	40.4	27.	2	2 44	F	244	7.7	1.65	2.62	1.36		2.96	1.09	4.52									1.98
ORF SEQ ID NO:	4.208A	20071		11215	11216	11361		12182	12183				13040		1300/2		130/3	11127				12081	12085			12931					14884	3 10099	4 10303
Exan SEQ ID NO:	7007	\$	8515	6179	6179	6312	6538	7069	2080	RORO		8982	8036	7000	3		8084	8098	7188	7287	8969	6977	6980	7197		7910		9220	9220	0686	0686	5113	5294
Probe SEQ ID 8	19	2848	3507	1177	1177	1315	150	2088	2088	90,	1000	3984	3019		3047		364	1001	2200	2312	1983	1992	1996	2220		2891	3921	4228	4226	4911	491	33	231

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Mus musculus a disintectin and metallicordehese demain (ADAM) 45 (metacristin) (Adams 42) and a	Potato Virus A RNA complete genome, ledate U	Mus musculus T-cell lymphoma invasion and metastasis 1 (Tram1) mRNA	Potato virus A RNA complete genome, Isolate U	Homo sapiens DKFZP686M0122 protein (DKFZP686M0122), mRNA	Homo sepiens DKFZP686M0122 protein (DKFZP588M0122), mRNA	Ovis aries prior protein gene, complete cds	Human papillomavirus type 7 genomic DNA	Fugu rubripes neurofibromatosis type 1 (NF1), A-tdnase anchor protein (AKAP84), BAW protein (BAW), and WSB1 protein (WSB1) genes, complete cds	Fugu rubripes neurofibromatosis type 1 (NF1), A-kinase anchor protein (AKAP84), BAW protein (BAW), and	WSB1 protein (WSB1) genes, complete cds	Homo sepiens Med4 homolog (MAD4) mRNA	602155687F1 NIH_MGC_83 Homo saplens cDNA clone IMAGE:4297556 5	601652250F1 NIH_MGC_82 Homo saplens cDNA clone IMAGE:3835556 5	M.mucedo gene encoding 4-Dihydromethyl-trisporate dehydrogenase	Cantharellus sp. partial 253 rRNA gene, isolate Tibet	Homo saplens putative pshHbA pseudogene for hair keratin, exons 2 to 7	Homo saplens zinc finger protein 157 (HZF22) (ZNF157) mRNA	Homo saplens zinc finger protein 157 (HZF22) (ZNF157) mRNA	Cotx lacryma-jobi dihydrodipicolinate synthase (dapA) gene, complete cds	Chlamydia mundarum, section 68 of 85 of the complete genome	601681233R1 NIH_MGC_72 Homo saplens cDNA clone IMAGE:3915945 3'	Figu rubripes gamma-aminobutyric acid receptor beta subunit gene, partial cds; 55kd erythrocyte membrane	protein (P55), synaptic vestcie-associated integral membrane protein (VAMP-1), procottagen C-proteinase	annancar protein (P.C.O.C.E.) genes, complete c>	22.2408.81 Sogres, 1648, jiver spicen_1NFLS_S1 Home sapiens cDNA clone IMAGE:431535 3	HISTIDINE-RICH PROTEIN PRECURSOR (GLONE PFHRP-III)	HISTIDINE-RICH PROTEIN PRECURSOR (CLONE PFHRP-III)	HISTIDINE-RICH PROTEIN PRECURSOR (CLONE PFHRP-III)	Homo saplans hypothetical protein PRO3077 (PRO3077), mRNA	Elaels oleifera sesquiterpene synthase mRNA, complete cds	pea seed-borne mosalo virus complete genome	
Top Hit Database Source	ţ	N-	7	N F	7	5	N	N.	Ę				EST HUMAN	EST HUMAN	N	NT	M	11	11	INT	NT	EST_HUMAN		!	Т	T		SWISSPROT	SWISSPROT	LN	NT.	NT	
Top Hit Acession No.	6752961 NT	131402.1	8678350 NT	131402.1		7661685 NT	167922.1		7			3		-		2.1		4507998 NT	4507998 NT		1.3E+00 AE002338.2					1.2E+00 AA6/0246.1				1.2E+00 8924234[h		252242.1	
Most Similar (Top) Hit BLAST E Value	1.5E+00	1.6E+00 AJ	1.5E+00	1.5E+00 AJ	1.4E+00	1.4E+00	1.4E+00 U67922.1	1.4E+00	1.4E+00	4 46400	1.45+00	1.45+00	1.4E+00 B	1.4E+00 B	1.3E+00 Z73640.1	1.3E+00 A	1.3E+00 Y19213.1	1.3E+00	1.3E+00	1.3E+00 U61730.2	1.3E+00	1.3E+00 B			1.35+00	1.2E+UU A	1.2E+00 P	1.2E+00 P05228	1.2E+00 P05228	1.2E+00	1.2E+00 A	1.2E+00 AJ	
Expression Signal	1.7	1.68	. 1.67	2.59	1.17	1.17	7.96	1.44	3.21	20	3.41	80.0	1.34	1.02	1.44	2.13	19.14	14.53	14.53	1.06	2.03	1.67		6	0.00	0.70	0.86	0.86	0.86	1.83	. 6.07	1.7	
ORF SEQ ID NO:		12438	12530	12438	10095	10096		12677	12771	49779	7/17	1		14944		10949		11318	11319					0	13330	1001	10867	10868	10869		11179	11224	
Exon SEQ ID NO:	5643	7318	7414	7318	5110	5110	7248	7580	7658	7858	32,52	3 60	9443	9968	2298	2909	6118	6277	6277	6336	6571	7448		C	0200	8	2834	5834	5834	5885	6147	6187	
Probe SEQ ID NO:	616	2344	2444	3065	30	30	2272	2598	2701	270.7	3250	277	20 20	4994	265	8	1112	1278	1278	1338	1574	2479		67.50		\$	813	813	813	867	1143	1186	

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	pes seed-borne motalic virus complete genoma	Homo saplens G-protein coupled receptor 14 (GPR14) gene, complete cds	Arabidopsis thallana DNA chromosome 4, contig treament No. 83	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 63	CONJUGAL TRANSFER PROTEIN TRBE PRECURSOR	Homo sapiens LHX3 gene, intron 2	Mus musculus subilisin-like serine protease LPC (PC7) gene, exons 1 to 9, partial cuts.	MR0-FT0175-050900-203-g06 1 FT0175 Homo septens cDNA	Homo sapiens LHX3 gene, intron 2	Rattus norvegicus Glycine receptor alpha 2 subunit (ghycine receptor, neonatal) (Glra2), mRNA	Rettus rattus cardiac AE3 gene, exons 1-23	Arabidopsis thallana DNA chromosome 4, contig fragment No. 21	Homo saplens post-synaplic density 95 (DLG4) gene, complete cds	T.pinnatum chloroplast rbcL gene, partial	Human mRNA for KIAA0227 gene, partial cds	QV0-BN0042-170300-163-g12 BN0042 Hamo saplens cDNA	Homa sapiens chromosome 21 segment HS21C013	Homo saplens chromosome 21 segment HS21C013	Homo sepiens hypothetical protein FLJ10749 (FLJ10749), mRNA	Homo sapiens hypothetical protein FLJ11280 (FLJ11280), mRNA	wf84h11.x1 Soarse_NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE:2359461 3' similer to SW:P631 HUMAN 012888 P53-BINDING PROTEIN 63RP1	Xylella fastidiosa, section 32 of 229 of the complete genome	Xylella fastidiosa, section 32 of 229 of the complete genome	Homo sapiens hypothetical protein FLJ10749 (FLJ10749), mRNA	Mus musculus protessome (prosome, mecropain) subunit, beta type 7 (Psmb7), mRNA	R.unicornis complete mitochondrial genome	African swine fever virus, complete genome	E.faecalis pbp5 gene	TELLURITE RESISTANCE PROTEIN TEHA	Mus musculus Konq1, Litpo5, Mash2, Tapa-1, Teso4 and Teso8 genes, alternative transcripts	Xenopus laevis rhodopsin gene, complete cds	Cavia cobaya mRNA for serine/threcine kinase, complete cds	Marchantia polymorpha genes for 26S rRNA, 5S rRNA, 18S rRNA, 5.8S rRNA and 26S rRNA
	Top Hit Database Source	F	F	Ę	N	SWISSPROT	N.	F	EST HUMAN	LN.	٦	Z.	N	L.	N	ĮZ.	EST_HUMAN	NT	TN	NT	ΝΤ	EST HUMAN	Ę	NT	Ž,	노	N	LN	LN T	SWISSPROT	NT	TN	NT	5
	Top Hit Acession No.	4J262242.1	4F140631.1				AF188740.1	J75902.1	3F373570.1	4F188740.1	6980951 NT	M87060.1	AL161509.2		Y09200.1	Γ		4L163213.2	4L163213.2	8922641 NT	8922973 NT	A1808360.1			8922641 NT	6755205 NT	5835331 NT	J18468.1	(78425.1	P25396	4,1251835.1			AB021684.1
	Most Similer (Top) Hit BLAST E Vatue	1.2E+00	1.2E+00	1.2E+00	1.2E+00		1.2E+00	1.2E+00	1.2E+00	1.2E+00	1.2E+00	1.2E+00	1.2E+00	1.2E+00	1.2E+00	1.1E+00	1.1E+00	1.1E+00	1.1E+00	1.1E+00	1.1E+00	1.1E+00	1.1E+00	1.1E+00	1.1E+00	1.1E+00	1.1E+00	1.1E+00	1.1E+00	_	1.1E+00	1.0E+00	1.0E+00	1.0E+00
	Expression Signal	1.7	76.0	6.13	6.13	3.29	0.73	8.37	2.04	76.0	1.01	2.1	0.99	1.92	5.44	1	1.36	7.24	7.24	0.72	2.43	0.93	1.32	1.32	0.95	0.88	8.1	4.28	1.11	0.71	0.97	3.83	2.17	2.22
	ORF SEQ ID NO:	11226	12048	13124	13125		13314	13641	13884	13314	14257			14405		10509			·	13443		13522		13648		13908		14796	14847	14968	14996		10194	
	SEQ ID NO:	6187	6946	8108	8108		8289		8881	8289	9266			9419							8487	8507	8642	8642	8842	8913	8088	9818	8478	9882	10028	2115	5185	5452
	Probe SEQ ID NO:	1186	1960	3092	3082	3213	3277	3630	3880	4174	4273	4346	4392	4429	4456	461	1725	3255	3255	3406	3479	3498	3636	3836	3840	3913	4094	4834	4899	5024	5058	86	113	415

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Table 4
Single Exon Probes Expressed in HBL100 Cells

		_	_	_	<u> </u>	T-	_	1				-	_	_	T	F	<b>&gt;</b>  [	<u>.                                    </u>	Ţ.		Щ	S	r	O	1	-	7	71	71	34	51
Ten Lil Pasadaka	mortiveed as do	Girardia tigrina mRNA for homeodomain transcription factor (so gene)	Homo saplens chromosome 21 segment HS21C018	Aedes aegypti mucin-like protein MUC1 mRNA, complete cds	V.carteri Algal-CAM mRNA	Plautie staff infestine virus RNA for nonstructural notwormtein general produit produit notwerness and		DNA GYRASE SUBUNIT B	3-OXO-5-41 PHA-STEROID 4-DEHYDROGENASE 1 (STEROID & AI PHA-STEROID 4-0 ETVER 1/ STEROID 4-0 ETVER 1/ STEROID 4-0 ETVER 1/ STEROID 8-0 E	3-OXO-5-ALPHA-STEROID 4-DEHYDROGENASE 1 (STEROID 5-AI PHA-BEN ICTASE 1/100 1/1	HYPOTHETICAL 67.9 KD PROTEIN CSF12.08C IN CHROMOSOMF I	ai 28g08 s1 Soares_lotal_fetus_Nb2HF8_9w Homo septens cDNA ctione IMAGE:1032830 3' similar to WP:C42D8.3 CE04204 :contains element MER22 MER22 repetitive element :	Xenopus laevis rhodopsin gene, complete cds	Agaricus bisporus mRNA for tyrosinase	Homo sepiens calclum channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced	Homo sapiens hypothetical protein FLJ10139 (FLJ10139), mRNA	Homo saplens chromosome 21 segment HS21C047	Taenia ovis 45W antigen (ToW4) gene, complete cds	Rattus norvegicus mRNA for N-acety/glucosaminytransferase III, complete ods	Pilot whale morbilityrus phosphoprotein (P) gene, partial cds	Human immunodeficiency virus type 1 proviral complete genome, Isolate 95MLB4	Homo sapiens chromosome 21 segment HS21C102	Apple mossic virus RNA 2 putative polymerase gene, complete cds	AMINO-ACID ACETYLTRANSFERASE (N-ACETYLGLUTAMATE SYNTHASE) (AGS) (NAGS)	Xenopus laevis rac GTPase mRNA, complete cds	Bromus inermis putative cytosolic phosphoglucomutase (pgm1) mRNA, complete cds	Bromus inermis putative cytosolic phosphoglucomutase (pgm1) mRNA, complete cds	PM2-UM0053-240300-005-f12 UM0053 Homo septens cDNA	Homo sapiens CGI-125 protein (LOC51003), mRNA	601675638F1 NIH_MGC_21 Hamo sepiens cDNA clane IMAGE:3868473 5	
Top Hit	Source	N	F	본	FX	<u> </u>	SWISSPROT	SWISSPROT	SWISSPROT	SWISSPROT	SWISSPROT	EST HUMAN	N.	N.	¥	N	LN LN	FN	ΤN	NT	NT	NT	NT	SWISSPROT	NT	NT	NT	EST_HUMAN	IN	EST HUMAN	
Top Hit Acession	oʻ	51660.1	63218.2	25984.1	416.1	1.0E+00 AB008531.1			. 800	800		53.1	1808.1	23816.1		1.0E+00 8922245 NT	83247.2	741.1		200817.1	45481.2	63302.2	9.9E-01 AF174585.1		9.8E-01 AF174644.1	97925.1	97925.1	799674.1	7705591	9.5E-01 BE902340.1	•
	BLAST E Value	1.0E+00 AJ2	1.0E+00 AL	1.0E+00 AF	1.0E+00 X80	1.0E+00/	1.0E+00 P48355	1.0E+00 P48	1.0E+00 P24	1.0E+00 P24	1.0E+00 O14228	1.00+000	1.0E+00 U23	1.0E+00 AJ2	1.0E+00	1.0E+00	1.0E+00 AL1	1.0E+00 l	1.0E+00 D10852.1	1.0E+00/	1.0E+00 AJ2	9.9E-01 AL1	9.9E-01	9.8E-01 P22567	9.8E-01	9.6E-01 AF	9.6E-01 AF	9.6E-01 AW	9.5E-01	9.5E-01	
5	angis	1.57	9.1	0.89	3.02	1.39	1.05	1.05	3.89	3.89	0.98	0.82	0.92	1.71	-	1.07	1.73	1.35	0.83	1.12	1.36	0.95	0.82	26.0	96.0	99.0	99'0	1.34	1.44	2.08	
ORF SEQ	<u>5</u>	10802	10703			11791	12504	12505	12841	12842	-	13163		13612	13943		14633					12647		10557		14291	14292	14312	12495	13699	
Exon SEQ ID	Ö	5604	5694	5695	7746	6714	7385	7385	7826	7826	7916	8142	5175	8604	8954	9148	9845	9736	9840	8954	10002	7529	8531	5554	7886	9307	9307	9327	7375	8698	
Probe SEQ ID	SO.	179	699	670	1365	1719	2414	2414	2806	2806	2897	3126	3519	2698	3956	4153	4880	4751	4829	4978	5031	5266	3524	519	2729	4315	4315	4336	2404	3694	

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Olingia Exteri Probas Expressed in ABL 100 Calls	Top Hit Descriptor	801675639F1 NIH_MGC_21 Hamo saplens cDNA clane IMAGE:3958473 5	Bartonella claridgelae RNA polymerase bela subunit (nodi) gene, partial cds	Pimpinella brachycarpa zinc finger protein (ZPP1) mRNA, complete cds	Homo saplens phylanoyl-CoA hydroxylase (PHYH) gene, exon 5	RC5-BT0503-271199-011-B01 BT0503 Homo sapiens cDNA	601441338T1 NIH_MGC_72 Home sapiens cDNA clone IMAGE:3918184 3'	Homo saplens hypothetical protein FLJ20048 (FLJ20048), mRNA	AB200GBR Infant brain, LLNL array of Dr. M. Soares 1NIB Homo sepiens cDNA clone LLAB200QB 5'	AB200G8R Infant brain, LLNL array of Dr. M. Soares 1NIB Homo septiens cDNA clone LLAB200G8 5'	Homo saplens neuredn III-alpha gene, perital cds	PUTATIVE FA20-DEPENDENT NADP REDUCTASE	Homo sapiens SOS1 (SOS1) gene, partial cds	nn05f11.s1 NCI_CGAP_Pr4.1 Hamo sapiens cDNA clane IMAGE:1078877	Pseudomonas aeruginosa topolsomerasa (top), pulative transcriptional regulatory protein OhbR (ohbR), ortho-	halobenzoate 1,2-dioxygenase beta-ISP protein OhbA (ohbA), OhbC (ohbC), ortho-halobenzoate 1,2- dioxonasea ainha-ISP protein OhhB (ohbB) and mits	Ret IGFII cene for insulin-like crowth factor II	244e03.r1 Soares [eta] heart NbHH19W Homo saplens cDNA clone IMAGE:343516 5	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 65	Drosophila melanogaster merlin (Dmerlin) mRNA, complete cds	Thermus thermophilus cytochrome c-552 (cycA) and CycB (cycB) genes, complete ods	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 18	Streptomyces antibioticus polyketide biosynthetic gene cluster	Rattus norvegious mRNA for RPHO-1, complete cds	IL3-CT0219-161199-031-C08 CT0219 Homo sapiens cDNA	Tanystylum orbiculare elongation factor 1-alpha mRNA, partial cds	Rettus norvegicus mRNA for RPHO-1, complete cds	Mus musculus TANK binding kinasa TBK1 (Tbk1) mRNA, complete cds	Homo sapiens MHC class 1 region	Homo sapiens MHC class 1 region	Staphylococcus aureus partial pia gene for phosphate actyltransferase allele 15	Bos faurus futb and ruff genes
Second Hox	Top Hit Database Source	EST HUMAN	N.	IN	F	EST_HUMAN	EST_HUMAN	LN	EST_HUMAN	EST HUMAN	N	SWISSPROT	L	EST_HUMAN	<u>,                                    </u>	<u></u> 5	Ę	EST HUMAN	NT.		NT	NT	NT	NT	EST HUMAN	NT	TN	NT	NT	NT	N N	Ę
Siligie	Top Hit Acessian No.	BE902340.1	9.4E-01 AF165990.1	Γ		BE071172.1		920	T26418.1	9.1E-01 T26418.1	Γ	Γ		8.7E-01 AA595863.1			Ī	Γ		8.6E-01 U49724.1		.2		8.2E-01 AB000489.1	AW376990.1	8.2E-01 AF063417.1	8.2E-01 AB000489.1		8.1E-01 AF055086.1			8.0E-01 AJ132772.1
	Most Similar (Top) Hit BLAST E Value	9.5E-01 B	9.4E-01	9.4E-01	9.3E-01	9.3E-01	9.2E-01	9.1E-01	9.1E-01	9.1E-01	9.0E-01	8.8E-01	8.7E-01	8.7E-01		8 7F.01	8 6E-01	8.6E-01	8.6E-01	8.6E-01	8.3E-01	8.3E-01	8.3E-01	8.2E-01	8.2E-01	8.2E-01	8.2E-01	8.1E-01	8.1E-01	8.1E-01	8.0E-01	8.0E-01
	Expression Signal	2.08	3.87	1.76	1.02	2.29	3.17	2.24	0.83	0.83	1.01	2.16	1.83	5.88			1.63	8.21	0.72	1.03	2.07	3.04	2.61	1.66	1.22	0.93	0.99	0.93	2.84	2.84	2.17	8.83
-	ORF SEQ ID NO:	13700					13205		13168	13169	14232	14386	10510	12839				10908	13552	13716				12086		13828	14890		13404	13405		10356
	Exon SÉQ ID NO:	8698	8144	8162	6689	7526	8182	7047	8148	8148	9248	9401	5489	7824		9820	5507	5867	8545	8714	5755	8038	8905	6982	7570	8821	9912	7642	8384	8384	5240	5344
	Probe SEQ ID NO:	3694	3128	3146	1694	2562	3166	2065	3132	3132	4254	4411	462	2804		4847	471	848	3539	3710	732	3021	3905	1999	2608	3819	4935	2684	3376	3376	176	286

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Table 4
Single Exon Probes Expressed in HBL100 Cells

		_	_	Т	_	_	_	_	_	_	7	_	_	_	_	_	_	_	T	Τ-	PC	Ŧ	~	₩	F	Т	P	1	<del>,                                    </del>	H	766
	Top Hit Descriptor	802072473F1 NCI_CGAP_Bm67 Homo saplens cDNA clone IMAGE:4215091 6'	Salmiri boliviensis olfactory receptor (SBO27) gene, perflai cds	Mus musculus gene for oviductal giycoprotein, complete cds	Nelsaeria meningittdis serogroup A strain 22491 complete genome; segment 7/7	G.gallus mRNA for nicotinic scetylcholine receptor (nAChR) beta 3 subunit	Mus musculus myosin IXb (Myo9b), mRNA	Lymantria dispar nuclear polyhedrosis virus gene for DNA polymerase, complete ods	Ureaplasma urealyticum section 31 of 59 of the complete genome	Homo saplens mRNA for KIAA1452 protein, pertial cds	Oryclalagus cuniculus mRNA for mitsugumin29, complete cds	Danio rario Trp4-associated protein Tap1A (tap1A) mRNA, complete cds	Gallus gallus SOX8 transcription factor (SOX8) mRNA, complete cds	601192033F1 NIH MGC 7 Hamo saplens cDNA clane IMAGE:3535785 5	Mus musculus eribidin (Emb), mRNA	Mus musculus embigin (Emb), mRNA	HSC1KH041 normalized Infant brain cDNA Homo saplens cDNA clone c-1kh04	EST371637 MAGE resequences, MAGF Homo septens cDNA	Rattus novegicus transmembrane receptor Unc5H1 mRNA, complete cds	RC3-CT0254-130100-023-c02 CT0254 Homo sapiens cDNA	Lycoperation hirsutum ADP-giucose pyrophorylese large subunit (AGP-L1) mRNA, complete eds	Mus musculus mejor histocompatibility locus class il region: mejor histocompatibility protein class il alpha	chain (iAalpha) and major histocompatibility protein class II beta chain (iEbeta) genes, complete ods; butvonhilin-like (NGs) hutomohilin-lix	CITRATE SYNTHASE	Homo sapiens PRO1975 mRNA, complete cds	Cotumix cotumix japonica sub-species japonica beta-actin mRNA, partial cds	Coturnix coturnix japonica sub-species japonica bela-actin mRNA, partial cds	Homo saplens chromosome 21 segment HS21C101	Homo sapiens FRA3B common fragile region, diadenosine triphosphale hydrolase (FHIT) gene, exon 5	Rattus norvegicus Protein phosphatase 1, catalytic subunit, beta isoform (Ppp1cb), mRNA	In14b09.x1 NCI_CGAP_Bm25 Homo septens cDNA clone IMAGE:2167677 3' similar to contains Aku repetitive element; contains element mit repetitive element;
יאסווין ווטאי	Top Hit Database Source	EST_HUMAN	N.	N.	NT	N	N.	Ę	Z	NT	N	F	Z.	EST HUMAN	1	Ę	EST HUMAN	EST HUMAN	N	EST HUMAN	N		<u>\</u>	SWISSPROT	N.	F.	LN LN	L	— <b>Ļ</b>	LN	EST HUMAN
1 Digitic	Top Hit Acessian No.	F530962.1	8.0E-01 AF127897.1	8.0E-01 AB006193.1			7657352 NT	Γ	7.9E-01 AE002130.1		7.9E-01 AB004816.1		7.9E-01 AF228664.1		6763745	6753745 NT	43785.1	W959567.1	187305.1	AW753353.1	F184345.1		AF050157 1	33915	F118085.1	F199488.1	F199488.1	L163301.2	F020503.1	6981387 NT	7.4E-01 Al598146.1
	Most Similar (Top) Hit BLAST E Value	8.0E-01 B	8.0E-01	8.0E-01	8.0E-01	8.0E-01	8.0E-01	7.9E-01 D11476.1	7.9E-01	7.9E-01	7.9E-01	7.9E-01 A	7.9E-01	7.9E-01 B	7.8E-01	7.9E-01	7.8E-01 Z	7.8E-01	7.8E-01 L	7.8E-01	7.7E-01 A		7.75-01	7.7E-01	7.7E-01	7.7E-01 A	7.7E-01 A	7.5E-01	7.5E-01	7.5E-01	7.4E-01
	Expression Signal	1.42	1.22	1.18	1.19	7.03	1.12	1.55	0.79	14.05	8.75	1.63	2.83	0.79	1.35	1.35	2.29	2.82	0.79	1.51	4.43		181	1.56	3.71	3.89	3.89	1.38	ţ-	0.95	1.25
	ORF SEQ ID NO:		13034	13273		14377		10503			12302	12303	13472		14449	14450		12311	14537		10223			12714	13538	14255	14258		10610	15006	11147
	Exan SEQ ID NO:	6969		8252	8628							7180	8446	9170	9469	6946	5883	7190	1556	9850	5208		6739		8527	9265	9265	5544	5611	10039	6119
	Probe SEQ ID NO:	1984	3003	3239	3821	4402	4822	451	705	1568	2201	2202	3438	4176	4479	4479	865	2213	4563	4871	143		718	2640	3520	4272	4272	209	579	5070	1113

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Homo saplens mRNA for KIAA0534 protein, partial cds	Malva pusilla actin (Act1) mRNA, complete cds	Homo saplens chromosome 21 segment HS21C046	xp83d04.x1 NCI_CGAP_Ov40 Homo saplens cDNA clone IMAGE:2746961 3' similar to contains element MER35 MER36 mentitive element:	Borrelia burgdorferi (section 52 of 70) of the complete genome	Homo sapiens HT017 mRNA, complete cds	Cicer anetinum partial mRNA for putative UDP alycose	Rattus norvegicus initiation factor-2 kinase (elF-2a) mRNA, complete cds	N. tabacum NelF-4A13 mRNA	Gallus gallus gene for melanocortin 2-receptor, complete cds	Fowlpox virus, complete genome	Glardia intestinalis variant-specifio surface protein (vsp417-6) gene, vsp417-8/A-I affele, complete cds	Human mRNA for KIAA0309 gene, partial cds	602035589F1 NCI_CGAP_Bm64 Hamo sapiens cDNA clane IMAGE:4183222 5	Limesenteroldes gene for sucrose phosphorylase (EC 2.4.1.7)	Homo saplens transcription factor IGHM enhancer 3, JM11 protein, JM4 protein, JM5 protein, T54 protein, JM10 protein, A4 differentiation-dependent protein, biple LIM domain protein 6, and synaptophysin genes, consider ads and I have relating channel as	onpose out, and Introduce account onemied as	Homo saplens transcription factor IGHM enhancer 3, JM11 protein, JM4 protein, JM5 protein, T64 protein, JM10 protein, A4 differentiation-dependent protein, triple LIM domain protein 6, and synaptophysin genes, complete cds; and L-type calcium channel a>	NUCLEOSIDE TRIPHOSPHATASE I (NUCLEOSIDE TRIPHOSPHATE PHOSPHOHYDROLASE I) (NPH		Rana catesbelana mRNA for bulifrog skeletal muscle calcium release channel (nanodine receptor) alpha Isoform/RNR1), complete ents	Hamo sapiens partial TCF-4 gene for T-cell transcription factor-4, exons 15-16	Mus musculus otogelin (Otog), mRNA	Mus musculus ologelin (Otog), mRNA		Homo sapiens mRNA for KIAA0614 protein, partial cds	to seplens cDNA clone IMAGE:288708 3' similar to	
Top Hit Database Source				EST HUMAN										EST_HUMAN						SWISSPROT							EST_HUMAN	
-	E	불	Z	83	토	Ę	Ę	Z	Z	Z	Ξ	Ę	Ę	<u> </u>	IN	2	4	Ż		<u></u>	<u> </u>	호	도 요	Z	N	N		
Top Hit Acession No.	7.4E-01 AB011108.1	AF112638.1	7.4E-01 AL163246.2	AW270842.1	7.3E-01 AE001168.1	7.3E-01 AF226421.1	7.3E-01 AJ400861.1	7.2E-01 L.29281.1	7.2E-01 X79140.1	7.2E-01 AB009605.1	AF198100.1	7.2E-01 AF065808.1	AB002307.1	7.2E-01 BF338350.1	D90314.1	7 2F-01 AF198779 1	1967.6.1	AF198779.1		P33066	7 15-01 024070 4	7.1E-01 AJ270777.1	7305360 NT	7305360 NT	7.0E-01 AB014514.1	7.0E-01 AB014514.1	N62412.1	
Most Similar (Top) Hit BLAST E Vætue	7.4E-01	7.4E-01	7.4E-01	7.4E-01 A	7.3E-01	7.3E-01	7.3E-01	7.2E-01	7.2E-01	7.2E-01	7.2E-01	7.2E-01	7.2E-01	7.2E-01	7.2E-01	7.2E-01	1.0	7.2E-01 A		7.2E-01 P33066	7.15.01	7.1E-01	7.1E-01	7.1E-01	7.0E-01	7.0E-01	7.0E-01	
Expression Signal	76.0	78.0	8.61	1.07	0.84	4.94	0.92	2.09	3.43	1.25	1.4	2.78	96.0	2.44	3.41	137	1	1.37		0.78	80	14.39	3.49	3.49	2.58	. 2.58	1.03	
ORF SEQ ID NO:	12376	13655	14161	14960	14453	14536	14986		11997	12485	13023	13398	13545	13787	14597	14904	1001	14805		14929	10719	13020	14070	14071	11248	11249	12474	
Exan SEQ ID NO:	7257	8649	1216	9984		L	10017	5841			8011	8378	L		6096	9826	١	9328		9951	6707		0806	0806	6029	6029	7363	
Probe SEQ ID NO:	2281	3643	4184	5013	4483	4562	5046	821	1917	2392	2993	3370	3533	3780	4624	4949		4949		4975	88	2989	4086	4086	1209	1209	2381	

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	yz/3e07.st Sosres_mulitple_sclerosis_2Nb-tMSP Homo sepiens cDNA clone IMAGE:288708 3' similar to contains Alu recettive element:	Homo saplens chromosome 21 segment HS21C101	Candida albicans squalene epoxidase (CAERG1) gene, complete cds and translational regulator gene, partial cds	Candida albicans squalene epoddase (CAERG1) gene, complete cds and translational regulator gene, partial cds	nn28a09.s1 NCI CGAP Gas1 Homo septens cDNA clone IMAGE:1085178.31	Chlamydia murdarum, section 3 of 85 of the complete cenome	Giardia Intestinalis carbamate kinase gene, complete cds	Synechocystis sp. PCC3803 complete genome, 27/27, 3418852-3573470	Rat(hooded) protectin gene ; exxn III and flanks	Homo sapiens nuclear factor of kappa light polypeptide gene enhancer in B-cells 1 (NFKB1) gene, complete eds	Homo explens nuclear factor of kappa light potypeptide gene enhancer in B-cells 1 (NFKB1) gene, complete eds	হধ?g12.e1 Soares_total_fetus_Nb2HF8_Dw Homo saplens cDNA clone IMAGE:786310 3' struiter to contains element TAR1 repetitive element;	Drosophila melanogaster Mst85C gene, complete eds; NMDMC isoform (Nmdmc) gene, complete cds, alternatively spliced; and transcription factor (Relish) cene. complete cds. alternatively spliced;	Mus musculus Wiskott-Aldrich syndrome protein (Wasp), mRNA	S.tuberosum mRNA for glucose-8-phosphate dehydrogenase	Homo sapiens lens epithelium-derived growth factor gene, alternatively spliced, complete cds	Homo saplens sema domain, seven thrombospondin repeats (type 1 and type 1-like), transmembrane domain (TM) and short cytoplasmic domain, (semaphorin) 54 (SEMA5A) mRNA	Calbicans random DNA marker, 282bp	Human hereditary haamochromatosia region, histone 2A-like protein gene, hereditary haamochromatosis (HLA-H) gene, RoRet gene, and sodium phosphate transporter (NPT3) gene, complete cds	H.wigaris Na,K-ATPase alpha subunit mRNA, complete cds	H.vulgaris Na,K.ATPase alpha subunit mRNA, complete cds	Mus musculus gene for Tob2, complete cds	Hamo sapiens SPP2 gene for escreted phosphoprotein 24 precursor, exons 1-8
Top Hit Database Source	EST HUMAN	N L	- <del>-</del> <u>-</u> <u>-</u>	-LV	EST HUMAN	N N	M	N	N	Ŋ	IN	EST_HUMAN	Ā	5	N.	NT		NT	NT	IN	IN	N	F
Top Hit Acession No.	7.0E-01 N62412.1	2	69674.1	69674.1	A593530.1	Γ				6.7E-01 AF213884.1	6.7E-01 AF213884.1	6.7E-01 AA451884.1	AF186073.1	TN 0828290		6.6E-01 AF199339.1	4506880 NT			6.5E-01 M75140.1			
Most Similar (Top) Hit BLAST E Value	7.0E-01	7.0E-01 A	6.9E-01	6.9E-01 U	6.9E-01	6.9E-01	6.8E-01	6.8E-01	6.8E-01 J00762.1	6.7E-01	6.7E-01	6.7E-01	6.7E-01	6.7E-01	6.7E-01 X74421.1	6.6E-01	6.6E-01	6.6E-01 Y07669.1	8.6E-01 U91328.1	6.5E-01	6.5E-01	6.5E-01	8.5E-01
Expression Signal	1.03	1.81	. 15.83	15.83	1.72	1.56	2.14	1.2	1.39	24.4	21.87	1.15	. 58	4	0.7	1.28	0.93	3.48	1.05	1.67	1.67	4.84	6.18
ORF SEQ ID NO:	12475		11004	11005	11334	13184	10992		14413	10366	10396	12179	12198	12963		12700	13446	13587		10647			14137
Exan SEQ ID NO:		9868	5971	5971	6290		5959	7565	9429	5353	5389	7007	7763	7946	9322	7588	8418	8581	8880	6645	5845	8357	9154
Probe SEQ ID NO:	2381	4889	955	955	1291	3148	942	2603	4439	296	337	2086	2102	2927	4331	2628	3409	3574	3983	618	618	3348	4159

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Single Exon Probes Expressed in HBL100 Cells

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Single Exon Plobes Expressed in ABL100 Cells	Top Hit Descriptor	Phaseotus vulgaris ATPase gamma subunit mRNA, nuclear gene encoding mitochondrial protein, partial cds	Homo saplens hypothetical protein PRO1580 (PRO1580), mRNA	Drosophila melanogaster 8kd dynein light chain mRNA, complete cds	Pseudomonas fluorescens tryptophan halogenase (pmA) gene, complete cds	Mus musculus dystroglycan 1 (DAG1) gene, exons 1 and 2 and complete cds	Homo saplens mRNA for KIAA1607 protein, partial cds	M.musculus whn gene	M.musculus whn gene	HISTIDINE-RICH PROTEIN PRECURSOR (CLONE PFHRP-III)	Hasmophilus Influenzae Rd section 4 of 163 of the complete genome	Shigella flexneri multi-antibiotic resistance locus	Gallus gallus bone morphogenetic protein 1 (BMP1) mRNA, partial cds	Gallus gallus bone morphogenetic protein 1 (BMP1) mRNA, partial cds	Lycopersicon esculentum p68a gane, complete CDS	Mus musculus secreted acidic cysteine rich glycoprotein (Sparc), mRNA	Homo saplens solute carrier family 28 (sulfate transporter), member 2 (SLC28A2) mRNA	Homo sapiens DNA for amyloid precursor protein, complete cds	Homo sapiens adaptor-related protein complex 3, mu 2 subunit (CLA20), mRNA	Human respiratory syncytial virus strain CH93-53b attachment protein (G) gene, complete cds	Viral hemorrhagic septicemia virus N, P, M, G, Nv, L genes, French strain 07-71	Xenopus mRNA for desmin	Homo saplens Notch3 (NOTCH3) gene, exans 26, 27, and 28	Haemophilus Influenzae Rd section 16 of 163 of the complete genome	Homo sapiens chromosome 21 segment HS21C067	Homo saplens chromosome 21 segment HS21C067	Rattus norvegicus cenexin 2 mRNA, partial cds	SIM1 PROTEIN	801852474F1 NIH_MGC_56 Hamo sepiens cDNA dane IMAGE:4076131 5	Vigna radiata mRNA for proton pyrophosphatase, complete cds	Megaselia scalaris sex-lethal homolog (Megsxl) gene, partial cds, alternatively spliced products	hl84f10.x1 NCI_CGAP_Kd13 Hamo saplens cDNA clone IMAGE:30059953'	APOLIPOPROTEIN A-1V PRECURSOR (APO-AIV)	APOLIPOPROTEIN A-IV PRECURSOR (APO-AIV)
KON Probes	Top Hit Database Source	Ž	7	Ę	Z	Ę	K	N	N.	SWISSPROT	N	NT	N	NT	NT	Ę	5	N.	7	NT	IN	NT	NT	NT	NT	NT	NT	SWISSPROT	EST HUMAN	NT	NT	EST HUMAN	SWISSPROT	SWISSPROT
alignic .	Top Hit Acession No.	28021.1	8924057 NT	48848.1	-	6.4E-01 U48854.2	6.4E-01 AB046827.1	12488.1	6.4E-01 Y12488.1		32689.1	81136.1	8.3E-01 U75331.1	75331.1	Y17275.1	6878076 NT	4557538 NT	6.0E-01 D87675.1	5802999 NT		6.0E-01 AJ233396.1		.1			6.9E-01 AL163267.2	5.9E-01 AF162756.1	6.8E-01 P40472				183.1		
	Most Similar (Top) Hit BLASTE Value	8.5E-01 U	6.5E-01	8.4E-01 U	8.4E-01	6.4E-01	6.4E-01	8.4E-01	6.4E-01	6.3E-01 P05228	6.3E-01	6.3E-01 U	6.3E-01	8.3E-01 U	8.3E-01	6.1E-01	6.1E-01	6.0E-01	6.0E-01	6.0E-01	6.0E-01	6.0E-01	8.0E-01	5.9E-01	5.9E-01	5.9E-01	5.9E-01	5.8E-01	5.8E-01	5.8E-01	5.8E-01	5.8E-01	5.7E-01 P06727	5.7E-01 P06727
	Expression Signal	2.32	1.39	. 8.85	0.0	3.74	1.18	0.74	0.74	3.16	1.73	25.01	1.82	1.82	0.67	3.53	0.98	1.08	2.76	1.38	0.81	2.18	1.03	1.03	9.18	9.18	3	1.05	0.97	4.1	1.31	2.29	1.04	1.04
	ORF SEQ ID NO:	14840	15008	10321	12594	13402	13778	14334		10488		12183	12587	12588			14379	10535			13727	13914		11031		13234		11943	13880	14353			11518	11519
	Exan SEQ ID NO:		ı	5310			8774	9355	9355	5469	5566	7079	7471	7471	7968		9386	5527	1			8923	9058	9000	8212	8212	8094	6855	8877	9374			6460	6460
	Probe SEQ ID NO:	4894	5073	250	2511	3374	3771	4364	4364	431	831	2098	2503	2503	2949	2330	4405	491	657	1345	3725	3923	4064	985	3196	3196	4100	1866	3876	4383	4652	4775	1463	1463

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	PUTATIVE TRANSCRIPTION FACTOR OVO-LIKE 1 (MOVO1) (MOVO1A)	Populus euramericana paaca-2 mSNA for 1-aminous/processes-1-ast-ondets austhose complete ad-	Droscohila extra sex combs dane, exon 1-4, complete cis	Homo saplens mRNA for KIAA0740 protein, pertial cds	Homo saplens mRNA for KIAA0740 protein, partial cds	Chicken TBP gene, exon 8, complete cds	Rattus norvegicus Proplany Coenzyme A carboxylase, beta palvaepiida (Poob), mRNA	GAG POLYPROTEIN (CONTAINS: INNER COAT PROTEIN P12, CORE PROTEIN P16, CORE SHELL PROTEIN P30: NUCLEOPROTEIN P101	GAG POLYPROTEIN (CONTAINS: INNER COAT PROTEIN P12, CORE PROTEIN P16; CORE SHELL PROTEIN P30: NITS FORROTEIN P30	Homo saplens superkiller viraitchic activity 2 (S. cerewisiae homolog). Ita (SKIV91.) mRNA	yo18a10.s1 Soares adult brain N2b5HB55Y Hamo sapians cDNA clone IMAGE:1782863'	Rabbit oral papillomavirus, complete genome	FOS-RELATED ANTIGEN-1	Bos taurus MHC class II beta-chain BoLA-DIB1 gane, partial cds	Homo sapiens KIAA0929 protein Max2 Interacting nuclear target (MINT) homolog (KIAA0929), mRNA	Homo sapiens KIAA0929 protein Max2 Interacting nuclear target (MINT) homolog (KIAA0929), mRNA	Pseudomonas syningae pv. tomato strain DC3000 AvrE (avrE), HrpW (hrpW), and GstA (gstA) genes, complete cds; and unknown genes	Pseudomonas syringae pv. tomato strain DC3000 AvrE (avrE), HrpW (hrpW), and GetA (getA) genes, complete cds: and unknown cenes	QV4-NN0040-070400-160-c04 NN0040 Hamo septens cDNA	Chlamydophila pneumoniae AR39, section 74 of 84 of the complete genome	Drosophila melanogaster mRNA for 15,15' beta carotene dioxygenase (beta-diox gene)	Homo saplens HLA class III region containing tenascin X (tenascin-X) gene, partial cds; cytochrome P450 21-Inydroxylase (CYP21B), complement component C4 (C4B) G11, helicase (SKI2W), RD, complement factor B (Bf), and complement component C2 (C2) genes,>	Homo saplens protein tyrosine phosphatase, receptor-type, zeta potypeptide 1 (PTPRZ1) mRNA	Homo sapiens protein tyrosine phosphatase, receptor-type, zeta potypeptide 1 (PTPRZ1) mRNA	Homo sapiens secreted C-type lectin precursor (LSLCL.) gene, complete ods
Top Hit Database Source	SWISSPROT							SWISSPROT	SWISSPROT		EST HUMAN		SWISSPROT					•	EST HUMAN			,			
	8	눌	ż	벌	ż	뉟	E	NS S	3	E	ES	ż	S	Ż	호	눌	돧	Ę	ES:	Z	Ż	Ę	Ż	3 NT	보
Top Hit Acession No.	5.7E-01 Q9WŢJ2	6.7E-01 AB033503,1	L41867.1	5.6E-01 AB018283.2	AB018283.2	D83135.1	8393912 NT	P03341	P03344	5902085 NT	6219.1	5.5E-01 AF227240.1	P48755	5.5E-01 U69097.1	7657286 NT	7657286 NT	AF232006.1	AF232008.1	5.4E-01 AW896087.1	5.4E-01 AE002247.2	AJ276682.1	AF019413.1	4506328 NT	450632B NT	AF087658.1
Most Similar (Top) Hit BLAST E Value	5.7E-01	5.7E-01	6.7E-01	5.6E-01	5.6E-01	5.6E-01	5.5E-01	5.5E-01 P03341	6.6E-01 Po	5.5E-01	5.5E-01 H4	5.5E-01	5.5E-01	5.5E-01	5.4E-01	5.4E-01	5.4E-01 AI	5.4E-01 A	5.4E-01	6.4E-01	6.4E-01	5.3E-01	6.3E-01	6.3E-01	5.3E-01 A
Expression Signal	1.38	2.37	96.0	1.11	1.11	2.28	1.79	2.21	221	0.79	1.33	3.6	1.37	66.0	13.58	13.58	1.11	1.17	2.4	2.35	1.53	1.61	10.07	10.07	3.67
ORF SEQ ID NO:	13188		14982	13320	13321	14095		12701	12702					14935	10224	10226	10811	10812			12293	10552	12779	12780	
Econ SEQ ID NO:	8168	8433	10013		8295	9111	6193	7589	7589		8010			9957	6210	5210	5612	5812			7172	6547			8185
Probe SEQ ID NO:	3152	3426	5042	3283	3283	4117	1192	2629	2629	2848	2992	3163	3608	4982	144	144	680	580	1252	2048	2193	612	2710	2710	3169

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Table 4
Single Exon Probes Expressed in HBL100

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	Top Hit Descriptor	Mycoplasma gentiatium section 9 of 51 of the campiste genome	Drosophila malanogaster helix-toop-helix mRNA, complete cds	NUCLEAR FACTOR OF ACTIVATED T CELLS 8 (T CELL TRANSCRIPTION FACTOR NEATS) (NF-ATS)	Homo septems phospholipid scramblase 1 dene, complete eds	Homo saplens chromosome 21 segment HS21C085	Homo sapiens mRNA for KIAA0740 protein, partial cds	Chlamydophila abortus strain S26/3 POMP91A and POMP90A precursor, genes, complete eds	Botrytis cherea strain T4 cDNA library under conditions of nitrogen deprivation	am77g05.s1 Stratagene schizo brain S11 Homo saplens cDNA clone IMAGE:1816504 S'	Medicago sativa chioroplast malate dehydrogenase precursor (p1mdh) mRNA, nuclear gene encoding chioroplast protein complete cits.	Avian infectious bronchitis virus isolate variant 2S1 spike alvoordein gene, partial cds	Homo saplens chromosome 21 segment HS21C081	Human adrenodowin reductase gene, exons 3 to 12	Polyanglum vitallinum (strain PI vt1) 165 rRNA gene	Polyangium vitellinum (strain PI vt1) 16S rRNA gene	w/39b12.x1 NCI_CGAP_Ut1 Homo sapiens cDNA done IMAGE:2427283 3'	TRANSCRIPTION-REPAIR COUPLING FACTOR (TRCF)	Homo saplens postmelotic segregation increased 2-like 9 (PMSZL9), mRNA	Homo saplens postmelotic segregation increased 2-like 9 (PMSZL9), mRNA	Buchnera aphidicola genomic fragment containing (chaperone HspS0) groEL, DNA biosynthesis initiating protein (dnaA), ATP operon (atpCDGAHFEB), and putative chromosome replication protein (ddA) genes.	complete cds; and termination factor Rho (rho) gene>	Buchnera aphidicola genomic fragment containing (chaperone Hsp80) groEL, DNA biosynthesis initiating protein (dnaA), ATP operon (atpCDGAHFEB), and putative chromosome replication protein (gidA) genes,	complete cds; and termination factor Rho (rho) gene>	Rattus norvegicus jagged protein mRNA, complete cds	Homo sapiens mRNA for KIAA1184 protein, partial cds	602076649F1 NIH_MGC_62 Homo saplens cDNA clone IMAGE:4243860 5	Xenopus laevis mRNA for c∽Jun protein, 1978 BP	Cavia porcellus pulmonary surfactant protein A (SP-a) mRNA, complete cds	oB2s08.s1 Soarss_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1626144 3*	802081103F1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:4245481 5'
	Top Hit Detabase Source	M	F	TORGREIME	LN DOWN	F	TN	NT	TN	EST_HUMAN	Ę	Ę	Ę	F	N.	N	EST_HUMAN	SWISSPROT	π	IT.		NT		NT	M	NT	EST_HUMAN	NT	N	- 1	EST_HUMAN
۰ŀ	<u> </u>	Z	Z	ď		Z	Z	Z	Z	<u>w</u>	2		Z	Z	Z	Z	Ш	S	4885552 NT	488552 NT		Z		Z	Z	Z	Ш	Z	Z	۳	=
	Top Hit Acesslan No.	U39687.1	1.20770.1	5 2F-01 O9WV30	6.2E-01 AF224492.1	5.2E-01 AL163285.2	6.2E-01 AB018283.2	U65942.1	5.2E-01 AL116780.1	5.2E-01 AA984165.1	AF020269 1	AF093796.1	5.2E-01 AL163281.2	M58509.1	5.1E-01 AJ233944.1	5.1E-01 AJZ33944.1	5.1E-01 Al858495.1	P96380	488	488		AF008210.1		AF008210.1	L38483.1	5.0E-01 AB033010.1	BF571482.1		U40869.1	4.8E-01 AA912842.1	BF693300.1
	Most Similar (Top) Hit BLAST E Value	5.3E-01	5.2E-01	5.2F-01	5.2E-01	5.2E-01	6.25-01	6.2E-01	5.2E-01	5.2E-01	5.2E-01	5.2E-01 A	6.2E-01	5.1E-01 N	6.1E-01	5.1E-01	5.1E-01	5.1E-01 P96380	5.0E-01	5.0E-01		5.0E-01 A		6.0E-01	6.0E-01	5.0E-01	4.9E-01	4.9E-01	4.9E-01	4.8E-01	4.6E-01 B
	Expression Signal	1.22	12.62	11.32	6.03	4.17	1.62	1.72	11.56	2.52	0.79	1.48	1.01	1.9	3.68	3.68	4.37	2.47	1.03	1.03		1.08		1.08	1.11	2.97	2.67	4.93	1.9	1.14	1.62
	ORF SEQ ID NO:		10857	11182	11208		12178	13071		13387				10843	10871	10672	13951	14043	12167	12168		12174		12175	13744	13788		11682	11941	-	13669
	Exan SEQ ID NO:	9084	5827	9150	8174	6836	7066	8063	8331	8368	8554	9349	9937	5640	9999	5668	8961	8056	7058	7058		7064		7084	8744	8785	5802	6616	6853	8468	8664
-		4090	908	1146	1171	1847	2085	3046	3321	3360	3547	4368	4960	613	640	640	3963	4062	2076	2078		2083		2083	3740	3782	781	1619	1864	3480	3658

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	602081103F1 NIH MGC_81 Hamo sepiens cDNA done IMAGE:4245481 6	455-00 at Soares fetal liver splean 1NFLS St Homo septens cDNA clone IMAGE:4041/8 3	DA SEAMENT AND RANGE SPECIFIC HEPARAN SULFATE PROTEOGLYCAN CORE PROTEIN	PRECURSOR (HSPG) (PERIECAN) (PLC)	Mus musculus DNA polymerase epsiron caralyas subum (r ors) years, consequences	COLLAGEN ALPHA 5(IV) CHAIN	ho60g02.x1 Soares_NFL_1 GBC_S1 name sapers conv. cons instance.co.	601657225R1 NIH MGC 6/ Hand septemble down with many many many many many many many many	Mus musculus minglial manipus and the precipitation of the precipitation (VEGF-B) (VEGF-B) (VEGF-B)	FACTOR)	Kattus nowegicus Oylickir-ti IIII vo., conspicus cas	Rattus nonegicus Symbol - Intrutto, Compress Constitution	7/91402.71 NCI_CCAP_ETID FINITE SEPTEMBLE CONT.	601237138F1 NIH _MGC_44 nailo sepais como cara image	Buzura suppressaria nucleopolyhedrosisvirus ecdysteroid UDP-glucosyltransferasa (egt) gene, complete ods	Callinrix lecchus MWAW opsin gens, upstream flanking region	Calithrix lacchus MW/LW opsin gene, upstream flanking region	CM2-DT0003-010200-077-c01 DT0003 Homo septens cDNA	MR0-BN0070-270300-008-g04 BN0070 Home saplens cDNA	Human somatostatin I gene and flanks	Califhrix lacchus MW/LW opsin gene, upstream flanking region	Callithrix lacchus MW/LW opsin gene, upstream flanking region	Xestia e-njarum granulovirus, complete genome	CELL DIVISION PROTEIN FTSH HOMOLOG PRECURSOR	Vivalle fastidinses section 93 of 229 of the complete genome	Ayrona users NHHMPu S1 Homo saplens cDNA clone IMAGE:1878945 3'	TABLET felal brain cDNA Homo septens cDNA clone 788IE14K similar to R07879, Z40498	ninghot st NCI CGAP Prt0 Homo sapiens cDNA clone IMAGE:997777 similar to gb:M33800 HLA CLASS	II HISTOCOMPATIBILITY ANTIGEN, DR-1 BETA CHAIN (HUMAN);	V77e01.r1 Soares Infant brain 1NIB Homo sapiens CUINA CIOIR INFORMATION	RC-BT091-210199-142 BT091 Homo saplens cDNA	AV705243 ADB Homo saplens cDNA clone ADBAHF08 5	
	Top Hit Database Source	EST HUMAN	Т	ESI HOMAN	ISSPROT	1	SWISSPROT	<b>EST HUMAN</b>	EST HUMAN	<u> </u>	SWISSPROT	Z	Į.	EST HUMAN	EST HUMAN	. 5	Ę	2 12	144441	EST HUMAN	ביים ביים	2 1	E LY	L	TOGGGGGGG	SWISSPACE	N	EST HOMAN	אייייטער ו פש	EST_HUMAN	<b>EST HUMAN</b>	FST HUMAN	FST HUMAN	
	Top Hit Acession No.	4 8E-04 RE693300 1 E	Ī	4.6E-01 AA677086.1		4F126378.1 NT		95.1		6680503 NT					4.4E-01 BE378707.1	4 45 04 1181154 1	T		T	T	AW8984//.1	J00306.1	AF155218.1	AF 1002 10.1		$\sim$ 1	~,	_	N81203.1	AA534083.1	_	-		
-	Most Similar (Top) Hit BLAST E Value	1 RE 04	4.01-01	4.6E-01	4.6E-01 Q06783	4.5E-01	4.5E-01 Q28247	4.5E-01	4.5E-01	4.4E-01	4.4E-01 P4	4.4E-01	4.4E-01	4.4E-01	4.4E-01	4 46 04	4.45	١	١										4.2E-01	4 2F.01				4.15-01
	Expression Signal	60	7.07	4.48	. 4.25	1 39	1.06	5.38	1.55	2.19	4.73	1.19	1.19	1.81	1.55					1.34					1.08	1.24	3.83		0.86	R 48				2.09
	ORF SEQ ID NO:	0000	136/0	12836	13277	43320	97001		14754		12420					_			10466		13018	14011	5 10465	10468	_	11389	13541		7					11122
	SEQ ID		8664	7819	8258	2000	1			L	7299	1	L			上	9915		3 5445		8004	9024	5445	2 5445			L			<u> </u>		16 9621		36 6093
	Probe SEQ ID NO:		3658	2789	2243	2470	3283	7180	4786	1985	23.5	3241	3241	3248	4113		4838	408	89	2801	2986	4028	4282	4282	4944	1341	3528	3552	3629		4558	4636	1077	1086

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	AV705243 ADB Homo septens cDNA clone ADBAHF08 6	PM-BT103-270499-884 BT103 Home sapiens cDNA	Homo saplens anaphase-promoting complex subunit 7 (APC7), mRNA	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 36	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 36	oj94b08.s1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1505943 31	Rhodococcus sp. AD45 isoG, isoH, Isol, isoA, isoB, isoC, isoD, isoE and isoF genes	cm33d02.s1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:1542619 3'	AV747880 NPC Hamo sapiens cDNA clane NPCBDF10 5'	Laqueus rubelius mitochondrion, complete genome	Drosophila melanogastar Dalmatian (dmt) mKNA, complete cds	Mus musculus platelet derived growth factor receptor, beta polypeptide (Pagins), mRNA	Mus musculus ubiquitin-protein ligase e3 componen n-recognin (Ubr1), mrtnA	Homo sapiens chromosome 21 segment HS21C080	Homo sapiens chromosome 21 segment HS21C080	Streptococcus pneumonies YilC (yilC), YilD (yilD), peniciliin-binding protein 2x (pbp2x), and undecaprenty-	prospitate-un-minnacipantepoure prospination and prospination of participates and construction of p	Ovis gries partial JD2 gene for T cell receptor delta chain (TCRDJ2), excn 1	Ovis aries partial JD2 gene for T cell receptor delta chain (TCRDJ2), exon 1	NADH-PLASTOQUINONE OXIDOREDUCTASE CHAIN 5, CHLOROPLAST	Gorilla gorilla carboxyl-ester lipase (CEL) gene, complete cds	Homo sapiens mRNA for KIAA1193 protein, partial cds	H. saplens B-myb gene	H.saplens B-myb gene	Sinorhizobium melijoti egi, syrb2, cys3 genes and ons	761d01.XI NC_CGAP_Brief home expense come invaces. 3538 for 3	Homo sapians protein Idnase PKNbeta (pknbeta), mKNA	Mus musculus pcm-1 mRNA for pericentriolar material-1, complete cos	Arabidopsis thaliana putative c-myb-like transcription ractor (MT 53R-3) mXNA, compress one	Mus musculus solute carrier tamily 1, member o (Skotac), minuka	Human immunodeficiency virus type 1 complete genome (1soute eo.cmr. 12.15)	Pieuronectes americanus aminopepusasse N (amply) gene, parual cus	Arabidopsis maliana Diva cilidinosonio 4, cong inginen ves co
Top Hit Database Source	EST_HUMAN	EST_HUMAN	T	NT.	N-	EST_HUMAN	NT	EST HUMAN	EST_HUMAN	_ _	님	- L	- 1	NT	NT			Ę	LZ.	SWISSPROT	NT	NT	NT	본	NT.	EST HUMAN	NT	IN	NT	NT.	LN L	Ę	¥
Top Hit Acession No.	05243.1	15949.1	7705283 NT	181536.2	161536.2	906344.1		309257.1	747880.1	4656	4.0E-01 AF203478.1	6679258 NT	6678490 NT	63280.2	63280.2		A DE-D1 AFORBODS 1		277511.1		18.1	_				59261	7019488 NT	3.8E-01 AB029291.1	3.8E-01 AF214117.1	6678002 NT	3.8E-01 AJ251057.1		4L161518.2
Most Similar (Top) Hit BLAST E Value	4.1E-01 AV7	4.1E-01 AIR	4.1E-01	4.1E-01 AL	4.1E-01 AL	4.1E-01 A	4.1E-01 AJ249207.1	4.1E-01 A	4.1E-01 AV	4.0E-01	4.0E-01 A	4.0E-01	4.0E-01	4.0E-01 AL1	4.0E-01		A DE-01	A PE OF A	4.0E-01 AJ	4.0E-01 Q31849	3.9E-01	3.9E-01	3.9E-01	3.9E-01 X82032.1	3.9E-01	3.9E-01 BF	3.8E-01		3.8E-01				3.8E-01 AI
Expression Signal	2.09	0.94	1.1	2.49	2.49	0.7	2.75	0.74	1.37	0.81	1.16	3.09	2.12	1.18	1.16		274	77.0	3.44	8.6	1.57	2.87	2.98	2.98	4.17	1.39	27.46	3.87	3.42	3.09	0.82	2	8.3
ORF SEQ ID NO:	11123	11632	12712	12908	12909				14509	11061	11365		10222	12937	12938		49830	floor			11407		12705	12706		13952			12578	12841	6		13438
SEQ ID NO:	6093	8570	7598	7888	7888	8244	9147	9172	9522	6031	6317	6451	5207	7917	L	L	0840	1		L			7593	7593	L	8962	5225	L		7775			8412
Probe SEQ ID NO:	1086	1673	2638	2869	2869	3228	4152	4178	4532	1022	1319	1454	2733	2898	2898			2012	3737	4672	1359	2571	2633	2633	3023	3964	159	502	2496	2560	2933	2974	3403

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Acession Top Hit Top Hit Descriptor Top Hit Descriptor Source	- #	W73812.X1 Soares NFL T GBC S1 Homo sapiens CLINA Gione IMAGE 2337633 3	PMA-HT0339-200400-010-G01 HT0339 Hamo sepiens CLINA	Homo sapiens mRNA for KIAA1410 protein, partial cds	Dania rerio bone morphogenetia protein 4 precursor (BMP4) gene, complete cas	EST21715 Adrenal gland tumor Homo sapiens cDNA 5' end	ok38c07.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Hamo sapiens cDNA clone IMAGE:1510188 3	MR3-0T0007-080300-104-b02 0T0007 Homo saplens cDNA	Nelsseria meningitidis serograup B strain MC58 section 50 of 206 of the complete genome	Human mibp gene, partial cds	yd03e05.r1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:24443 5	yd03e05.r1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMAGE:24443 6	hg33f02x1 NCI_CGAP_GC8 Hamo sepiens cDNA clone IMAGE:2947419 3'	hg33f02.x1 NCI_CGAP_GC6 Hamo saplens cDNA clone IMAGE:2947419 3'	Mus musculus ribosomal protein S19 (Rps19) gene, complete cds	Human mRNA for KIAA0323 gene, partial cds	P.Irregulare (P3804) gene for actin	RC5-ST0171-181099-011-007 ST0171 Homo septems cDNA	PROTEIN-L-ISOASPARTATE O-METHYLTRANSFERASE (PROTEIN-BETA-ASPARTATE	METHYLTRANSFERASE) (PIMT) (PROTEIN L-ISOASPARTYL METHYLTRANSFERASE) (L-  ISOASPARTYL PROTEIN CARBOXYL METHYLTRANSFERASE)	Drosophila malanogaster sugar transporter 3 (sut3) mRNA, complete cds	H.sapiens serotonin transporter gene, exons 9 and 10	H.saplens serotonin transporter gene, exons 9 and 10	RC1-HT0545-150600-014-b12 HT0545 Homo seplens cDNA	Z.mays mRNA for caseln kinase II alpha subunit	ha02g04.x1 NCI_CGAP_Lu24 Homo saplens cDNA clone IMAGE:2872568 3'	Treponema palitdum section 3 of 87 of the complete genome	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 36	Mus musculus mannose receptor, C type 2 (Mrc2), mRNA	Hano saplens GAP-like protein (LOC51306), mRNA	Homo sapiens GAP-like protein (LOC51306), mRNA	601811060R1 NIH_MGC_48 Homo sapiens cDNA clone IMAGE:4053951 3	801894653F2 NIH_MGC_19 Hamo sapiens cDNA clone IMAGE:4124244 6	
Top Hit Detebase Source		EST HUMAN	EST HUMAN	L		EST_HUMAN	EST_HUMAN	EST HUMAN			EST_HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	NT	NT	Į.	EST HUMAN		TORISSPECT	IN	¥	NT.	EST HUMAN	N	EST_HUMAN	FN	LN	11	[ <del>1</del> ]	15	EST_HUMAN	EST HUMAN	
Top Hit Acession No.		19.1	080.1	831.1	3.7E-01 AF056336.1 NT		1.70	8037.1	3.7E-01 AE002408.1 NT				3.6E-01 AW590184.1   E		3.6E-01 AF216207.1 N		Г	3.1			485.1	Γ		-	Γ	9393.1	AE001187.1	AL161536.2		7706138 NT	7708136 NT	3.5E-01 BF129796.1		
<u>≅</u> ≈ m	Value	3.8E-01/	3.8E-01 BE154	3.7E-01 AB037	3.7E-01	3.7E-01	3.7E-01 AI2187	3.7E-01 AW87	3.7E-01	3.6E-01 U89241.1	3.6E-01	3.6E-01 T80255.1	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01	3.6E-01 AW81		2 EC. 01 D2420	3.0E-01	3 6E-01	9 AE 04	3.8E-01	3.6E-01 Y1152	3.6E-01	3.6E-01	3.5E-01		3.6E-01	3.5E-01			
Expression Signal		0.77	0.93	5.9	10.03	1.12	7.57	1.55	3.38	8.26	2.38	2.38	3.81	3.81	4.6	78.0	2.62	0.95			7.88	183	1 83	88.0	1.49	2.79	1.25	0.85	2.22					
ORF SEQ ID NO:			13679	12489	13407	13786	14089	14167	14248		11339	11340	11949	11950	11893	L		12508			ROOT	13420			L	L			L					]
SEQ ID		8469	8676	7381	8386	8782	9102	9186	9258	5995	6293	6293	6861	6861	6898	7185	7300	7388			10055	L			L	L		L	1	L				
Probe SEQ ID		3461	3671	2410	3378	3778	4108	4183	4265	980	1295	1295	1872	1872	1912	2208	2326	2417			8007	2388		3368	4610	4840	5037	118	2 00	715	7,5		1580	

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no11b10.s1 NCI\_CGAP\_Phs1 Homo sapiens cDNA clone IMAGE:1100347 3'
MRA-BT0403-230200-202-c01 BT0403 Homo sapiens cDNA
qj95c05.x1 NCI\_CGAP\_Kid3 Homo sapiens cDNA clone IMAGE:1867208 3' similar to contains Alu repetitive Canis familiaris rod photoreceptor cGMP-gated channel alpha-subunit (CNGC1) mRNA, complete cds
Homo sepiens pulmonary surfactant protein D, promoter region and exon 1
Methylovorus sp. strain SS1 putative GrpE (grpE), DnaK (dnaK), and putative DnaJ (dnaJ) genes, complete Hamo saplens partial N-myc (exon 3), HPV45 L2, HPV45 L1, HPV46 E8, HPV45 E7 and HPV45 E1 genes π08e09.s1 Stratagene NT2 neuronal precursor 637230 Homo saplens cDNA clone IMAGE:650872 S¹ nr60d03.s1 NCI\_CGAP\_Lym3 Homo saplens cDNA clone IMAGE:1172357 S¹
Danio rario homeobox protein (hoxb5b) gene, complete cds 7n94e01.x1 NCI\_CGAP\_Ov18 Homo sepiens cDNA clone IMAGE:3572232 3' similar to TR:Q9UJ15 Rattus novegicus ADP-ribosylation factor-directed GTPesse activating protein mRNA, complete cds HOMEOBOX PROTEIN HOX-A4 (HOX-1.4) (MIH-3) 602184016T1 NIH\_MGC\_42 Homo sapiens cDNA clone IMAGE:4300261 3' Mus musculus distritegrin 6 (Dtgn5), mRNA Peeudomonas flucrescens cofR, cofS genes, orf222 and partial InaA gene Azobbacter vinelandii niA gene for NifA protein (positive regulatory element) Synechocystis sp. PCC8903 complete genome, 11/27, 1311235-1430418 Homo saplens chromosoms 21 segment HS21C010 Homo saplens chromosome 21 segment HS21C010 Synechocystis sp. PCC6803 complete genome, 11/27, 1311235-1430418 Arabidopsis thaliana DNA chromosome 4, contig fragment No. 45 Homo saplens KIAA1100 protein (KIAA1100), mRNA Rat leukocyte common antigen (L-CA) gene, exons 1 through 5 Top Hit Descriptor Rhizobium leguminosarum sym plasmid pRL5JI nodX gene Rhizobium leguminosarum sym plasmid pRL5JI nodX gene isolated from IC4 cervical carcinoma cell line PROLINE-RICH PROTEIN LAS17 Single Exon Probes Expressed in HBL100 Cells Q9UJ15 DJ18C9.1; ş EST HUMAN EST HUMAN EST\_HUMAN EST\_HUMAN NT EST\_HUMAN EST\_HUMAN SWISSPROT SWISSPROT Top Hit Detabase Source Ę 눋 눋 눋 Ę 눋 눌 뉟 뉟 7662485 Top Hit Acession 3.3E-01 X07990.1 3.3E-01 X07990.1 3.3E-01 AL161545.2 3.3E-01 3.4E-01 BF449010.1 3.4E-01 AA584198.1 3.4E-01 BE069912.1 BF568880.1 3.4E-01 AL163210.2 3.4E-01 AL163210.2 3.4E-01 AF108835.1 3.5E-01 AF071253.1 3.4E-01 AI240973.1 3.5E-01 AA223252.1 AA642138.1 3.4E-01 AJ242956.1 3.4E-01 U83905.1 3.4E-01 AF034862.1 ģ 012446 3.4E-01 Y09798.2 3.4E-01 Y00554.1 **L35776.1** 3.5E-01 M18349. 3.4E-01 D90909. D90909. 3.5E-01 P06798 3.3E-01 3.3E-01 3.5E-01 3.6E-01 (Top) Hit BLAST E Aost Simila 1.18 4.55 1.07 2.49 4.08 0.92 1.67 4.38 1.48 5.81 1.76 1.56 8.0 0.8 £. 1,8 2.06 5.2 Expression Signed 13119 10078 10078 10500 10860 11221 11331 11628 13488 14478 14118 12433 12968 12969 13106 14748 11010 13303 11657 12318 12616 11351 ORF SEQ ÖNQ 8705 9778 5655 6183 6287 6566 5094 9135 9761 6721 5976 8281 6304 8937 5094 6596 7774 8721 7951 **2** SEQ ID ğ 445 627 1288 1569 2531 4140 4777 3453 3839 4509 794 <del>1</del>90 961 1306 2338 2932 3088 3268 Probe SEO ID 22 22 23 80

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	Top Hit Descriptor	EST36722 Embryo, 8 week I Homo saplens cDNA 5' end	Methylococous capsulatus strain Bath outer membrane protein Mopb (mopb) gene, outliness cue	Homo sepiens undine monophosphate synthetese (orotate phosphoribosy transferase and crotdine-o-	decarboxylase) (UMPS) mRNA	Bacteriophage phi-YeO3-12 complete genome	INTERLEUKIN-12 ALPHA CHAIN PRECURSOR (IL-12A) (CT 10 LOAC LIMT NOCH LEMAN CICALLE). FACTOR 35 KD SUBUNIT) (CLMF P35)	Streptomyces argillaceus mithramycin blosynthetio genes	Homo sapiens MTA1-L1 gene, complete cds	EXODEOXYRIBONUCLEASE V BETA CHAIN	GENOME POLYPROTEIN (CONTAINS: N-TERMINAL PROTEIN (P1); HELPER COMPONENT	PROTEINASE (HCPRO); PROTEIN P3)	Arabidopsis thaliana DNA chromosome 4, contig magment no. 10	Hypoxykon fregiforme chitin synthese gene, perusa cus	Homo sapiens RAS protein activator like 1 (GAP1 like) (RASAL1) minnA	Rettus norvegicus DNA for regucalcin, partial cds	IB78b12-X1 NCI CGAP US Home septens cDNA done IMAGE:2205407 3' similar to go:X5/622 AN HOELY	PEPTIDE TRANSPORTER 1 (HUMAN);	Symetricoysus sp. 1 Cocco Company 1 NFIS S1 Homo saplens cDNA clone IMAGE: 1655392 3' stmilar to	contains element MER4 repetitive element;	Rattus novegicus EH domein binding protein Epsin mRNA, complete cds	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 61	Fusarium pose virus 1 RNA2 putative RNA dependent RNA polymerase gene, compress cus	P.wigaris arc5-1 gene	LACTOSE PERMEASE (LACTOSE-PROTON SYMPORT) (LACTOSE INVISED OF TROUCH)	S.cerevisiae chromosome II reading frame ORF YBR172o	EST369264 MAGE resequences, MAGD Homo septems cDNA	EST369264 MAGE resequences, MAGD Homo sapiens cDNA	601868804F1 NIH MGC_17 Homo sapiens cDNA clone IMAGE:4111512 6	Mus musculus Phylknotted 1 homeobox (Pknox1), mRNA	Home saniens promyelocytic leukemla zinc finger protein (PLZF) gene, complete cds	112-UT0073-180900-161-H11 UT0073 Hamo saplens cDNA	
	Top Hit Database Source	EST_HUMAN			7	NT	SWISSPROT	¥	5	SWISSPROT	_	SWISSPROT	NT	NT.	7	LZ.		EST HUMAN	LN	FST HUMAN	Ė	LN	LN	LZ	SWISSPROT	TN	EST HIMAN	EST LIMAN	EST HIMAN	TO LICE	12 2	EST HUMAN	
218:110	Top Hit Acession No.	3.3E-01 AA332734.1 E	F031148.1		4507834 NT	251805.1	27.48	Τ	Τ	3.3E-01 ABU12822.1	Τ		38.2		4759025INT	3 3F-01 D31682.1		3.3E-01 AI539114.1		A 1024002 4			Ī	50202.1	OABROA					BF20381		AF060568.1	
}	Most Similar (Top) Hit BLAST E Value	3.35-01/	3 3E-01		3.3E-01	3.3E-01 AJ	0.00	9.25.01	20.0	3.35-01	3.0E-01	3.3E-01 P22602	3.3E-01	3.3E-01	3.3F-01	3 3F-01		3.3E-01	3.3E-01									3.22-01				3.21-01	
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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Rabbit beta-like globin gene cluster encoding the epsilon, gamma, deta (pseudogene) and beta globin polypeptides, complete eds	HYPOTHETICAL 81.7 KD PROTEIN C13G7.04C IN CHROMOSOME I TRECORSON	602081972F1 NIH_MGC_81 Hamo sapiens CUNA dana IMAGE:4240300 0	Arabidopsis thaliana DNA chromosome 4, contig tragment No. 20	ye90h06.r1 Soeres fetal fiver spleen 1NFLS Home sapiens cunA done invade::123031 5 similar to gb:3/84241 QM PROTEIN (HUMAN);	Homo saplens KIAA0174 gene product (KIAA0174), mKNA	Homo sapiens KIAA0174 gene product (NAA0174), minaan	1148h08x1 Scares NRL I GBC S1 Home septents curve cigne invence. 281 0031 5	Mus musculus gene for Ser/ I fir Kinase Knikawike, excit o	Daticus carda mKNA for transcription ractor Ezer (Ezer gene)	Xylella fastidiosa, section 130 of 228 of the complete gamonie	Mus musculus protein kinase C, epsilon (PKce), mxivA	Homo saplens Xq pseudoautosomai region; segment 1/2	XX63(08.X1 NCI_CGAP_KIG11 Home sapiens cuive immoli	Basehoptera priyeatus gene entotuning atrea marando popula.	A.Immereus pussure gene encounty integrace, make v	Colynebacterum sp. ALT-1 alyn O gain or polygram commission.	Unest aminos Varseaudos thesometreologis segment 2/2	Home saptans Ay pseudoauceania region, egon, egonemente per en	Dalatalopida priyaata yano aroonig Racterionbane APSE-1, comblete genome	Anufex applicing section 68 of 109 of the complete genome	Xenous leavis transcription factor E2F mRNA, complete cds	PM1-CT0328-171299-001-f12 CT0326 Homo septens cDNA	DAM CT0328-171299-001-412 CT0328 Home sapiens cDNA	LONG COAP GREAT HOND Sapiens CDNA clone IMAGE:2188412 3' similar to gb:D16050 NIL-2-A	ZINC FINGER PROTEIN (HUMAN); contains element L1 repetitive element;	Cavia porceilus mRNA for giutamione a-ransterase, cumpreso cos	2557412.11 NCI_CGAP_GCB1 Hamp sapiens curve giors investigated to sering a commercial constitue plantant.	Home sanlens chromosome 21 segment HS21C007	
dy cogo! I love ellino	Top Hit Database Source		SWISSPROT	L HUMAN		T_HUMAN			EST_HUMAN						EST HUMAN				EST HUMAN					TOWAN	TOT TOWAR	EST HOMAN	EST_HUMAN	T		NAMOE I SIL	_
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olifile.	Top Hit Acession No.	118818.1	10268	3.2E-01 BF693617.1	L161514.2	118051.1		7881971 NT	\W629036.1	3.1E-01 AB029069.1	J251586.1	3.1E-01 AE003984.1	6755083 NT	J271735.1	3.0E-01 AW300400.1	3.0E-01 AJ006755.1	3.0E-01 X99082.1	3.0E-01 AB030481.1	3.0E-01 AW817785.1	3.0E-01 AJ271738.1	3.0E-01 AJ008755.1	3.0E-01 AF15/835.1	2.9E-01 AE000/36.1	2.9E-01 AFU/0111.1	2.9E-01 AW 754239.1	AW754239.1	AI610836.1	AB016426.1			AL163207.2
ŀ	Most Similar (Top) Hit BLASTE Value	3.2E-01 M1	3.2E-01 Q1	3.2E-01 B	3.2E-01 AL	3.1E-01 R	3.1E-01	3.1E-01	3.1E-01 AV	3.1E-01	3.1E-01 AJ	3.1E-01	3.0E-01	3.0E-01 A	3.0E-01	3.0E-01	3.0E-01	3.0E-01	3.0E-01	3.0E-01			2.9E-01	2.9E-01	2.9E-01	2.9E-01	2.9E-01				2.9E-01
	Expression Signal	£.	1.43	7.8	1.02	232	272	2.72	1.27	3.43	0.89	5.69	1.59	9.02	1.65	3.47	. 1.03	4.06	1.61	0.98	2.26	0.89				2.82	0.7	1			0.85
	ORF SEQ ID NO:	14253	14342		14994	12680	12696	12697			13820	14772					11837		13781	13871	14352					13217	13800			14329	4
	Exan SEQ ID NO:	8263	9362	9585	10025	7562	10077	2709	982	8116	8814	8790	7689	L	L	1		L	8777	8868	9373			8126	8194	8184	7080	L		9850	9534
!	Probe SEQ ID NO:	4270	4369	4597	5054	Sano	2626	2825	2785	3100	3811	4806	ž	25.2	1204	1478	1760	3139	3774	3867	4382	4589	1973	3110	3178	3178		200	ROAS	4359	4545

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	we08t03.x1 NCI_CGAP_Kid11 Homo sepiens cDNA done IMAGE:2287309 3' similar to contains L1.t2 L1 repetitive element;	Arabidopsis thallana DNA chromosome 4, contig fragment No. 81	Rattus norvegicus A-kinase anchoring protein AKAP150 mRNA, complete ods	Prune dwarf virus movement protein, complete cds; cost protein, complete cds	Guina guina cocyte maturation factor Mos (c-mos) gene, partial cds	601148733F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:3163688 6	601148733F1 NIH_MGC_19 Homo sepiens cDNA clone IMAGE:3163688 6"	Human mRNA for serine/threonine protein kinase, complete cds	QV1-CT0364-120200-065-b05 CT0364 Homo sepiens cDNA	DKFZp588i2321_r1 586 (synonym: hute1) Homo sapiens cDNA clone DKFZp588i2321	hd44b03.x1 Soares_NFL_T_GBC_S1 Homo saplens cDNA clone IMAGE:2912333 3'	Escherichia coli K-12 MG1655 section 384 of 400 of the complete genome	Escherichia coli K-12 MG1655 section 384 of 400 of the complete genome	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 65	Arabidopsis thaliana mRNA for lipoyltransferase, complete cds	Toxoplasma gondii 90kDa heat-shock protein (HSP90) mRNA, partial ods	B.taurus microsatellite (ETH121)	B. taurus microsatellite (ETH121)	Pyrococcus horikoshii OT3 genomic DNA, 777001-894000 nt. position (4/7)	Borrella burgdorferi (section 68 of 70) of the complete genome	Pseudomonas aeruginosa PA01, section 11 of 529 of the complete genome	ov44g10.xt Soares_lestis_NHT Homo sapiens cDNA clone iMAGE:1640226 3' similar to contains Alu repetitive element.contains element MER22 repetitive element;	RNA POLYMERASE BETA SUBUNIT (LARGE STRUCTURAL PROTEIN) (L PROTEIN)	Bovine adenovirus 3 complete genome	602042601F1 NCI_CGAP_Bm67 Hamo saplens aDNA alone IMAGE:4180129 6	q59c11.x1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:1876628 3' similar to contains Alu	repetitive element.contains element LTR6 repetitive element ;	oa41h01.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1307669 3	Rettus novegicus CDK104 mRNA	zx39b10.s1 Soares_tota_fetus_Nb2HF8_6w Homo sepiens cDNA clone IMAGE:788827 3' similar to contains Alu repetitive element;	ipomoea purpurea transposable element Tip100 gene for transposase, complete cds
Top Hit Database Source	EST_HUMAN	TN	LN	Ę	NT	EST_HUMAN	EST HUMAN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	IN	TN	NT	NT	NT	NT	NT	NT	NT	EST HUMAN	SWISSPROT	NT.	EST_HUMAN		EST HUMAN	EST_HUMAN	F	EST HUMAN	¥
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Most Similer (Top) Hit BLAST E Velue	2.9E-01 A	2.9E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01	2.8E-01 A	2.8E-01	2.8E-01		2.8E-01	2.8E-01		2.8E-01				2.8E-01			l							
Expression Signal	1.33	1.01	211	2.0	2.64	0.98	96'0	0.92	2.03	1.44	1	2.08	2.06	1.49	1.4	1.35	2.3	2.3			0.91	236		2.85			2.3	1.68	2.82		
ORF SEQ ID NO:					11108	11298	11299		11761	12050	12165	12483	12494		12871		12941	12942		13888	0		14313		١		14701	14950	10522		
Exam SEQ ID NO:	8923	1666	5597	5601	6075	6256	8256	6270	6685	6948	7055	7372	7372	7445	7657	7920	7921	7921	8305	L.	9001	auda	L	19867			9716		L		
Probe SEQ ID NO:	4946	6020	883	895	1067	1258	1258	1272	1689	1962	2073	2401	2401	2476	2594	2901	2902	2902	3294	3890	4005	4075	4337	4682	4711		4731	5004	473	8	1242

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	Top Hit Descriptor	Glamhlia SB2 seens	2222/10 of Source (atal heart NKHHstow Lymns and heart NKHstow	GAG POLYPROTEIN (CONTAINS: INNER COAT PROTEIN P12; CORE PROTEIN P16; CORE SHELL PROTEIN P36; MI I'M EADBATTEIN PAGE.	Rathe portraction to fail and the second sec	Feline Immi inclessions with a management of the contract of t	1843-11.72 NCI_CGAP_Lu25 Home septems cDNA clone IMAGE:2048838 3' straitar to contains element L1	CM4 LT0276 Defined age - of Hanseltt	Web2et1 4 NCL CoAB Kitat III	Home series Digests 2 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -	Tribon copyrists of the Chical Poglon, telomeric end	Muvain essuvim (W. Gsod) gene, complete cds	INCITATIONS A SECURITIONS ASSUMED SEPTEMBERS CONA	Bos frame - Palls S.	SOJEJOSSE VIII VOO TATI	Charles may a march 1 from sapiens a DNA clone IMAGE:3912345 6	Arehidoneis the line DNA -	Arabidosele theliana DNA at	bboddin Y NIH MCC 14 Home 2015	RECONSTRUCTION OF THE SEPTEMBERS CLINA CICHIE IMAGE: 2558461 3' SIMILIAT to gb: M36072 605 MACHINES COMPL. PROTEIN L7A (HUMAN); gb:M14689_cds1 Mouse surfet locus surfet 3 protein gene	Independent of the second of t	B. martimus that serve	80126016F1 NIH M.C. 9 Lime and 12 Line	EST386635 MAGE resecuences. MAGM Home content contents.	Appearance of the second of th	FST377580 MACE	OV1-BT0630-040400-132-473 BT0630 Home septems culva	Enterococcus faecium strain N97-330 vanD giycopepide resistance gene cluster, complete cds; and	BBSB007.11 Stratagement failed refiles 0372002 Homes continue and a file of the continue and a file of	Arabidopsis thaliana PSI type III chlorophyl a/b-binding gratein (Lhosa*1) mRNA complete A-de
2000	Top Hit Database Source	LN	EST HUMAN	SWISSPROT	Į.	Į.	- H	EST HIMAN	EST HIMAN	TN		EST LIMAN	SWISSPBOT	TN	EST HIMAN	A CANONIA PAR	TA L	5		TOUR TOUR	TN FIN	Ę	EST HUMAN	EST HUMAN	-	EST HIMAN	Т		HUMAN	
	Top Hit Acession No.	Γ			T	2.7E-01 Y13868.1		2.7E-01 BF088284 1		T	Τ	2.7E-01 AW858131 1	Τ	Т	885087 4	T	2.6E-01 AL161472.2	Г	Τ	2 6F-01 AW7934524	1844.1		Ţ.			Τ	2.6E-01 BE080598.1		Τ	
	Most Similar (Top) Hit BLAST E Value	2.7E-01	2.7E-01 W58087.1	2.7E-01	2.7E-01 /	2.7E-01	2.7E-01	2.7E-01	2.7E-01	2.7E-01 L77569.1	2.7E-01	2.7E-01 A	2.6E-01 P	2.6E-01.D1	2.6E-01 B	2.6E-01 A	2.6E-01	2.6E-01 A		2 6F-01 A	2.6E-01 M1	2.6E-01 Y12896.1	2.6E-01 B	2.6E-01 A	2 6F-01 A	2.6E-01 A	2.6E-01 BI	2.6E-01 AI	2.6E-01 A	2.6E-01 U01103.1
	Expression Signal	1.3	221	1.16	1.42	6.34	2.78	0.72	1.98	2.26	1.14	4.25	1.71	60.	1.36	0.92	4.92	4.92			2.5	3.6	6.05	1.09	2.15	0.94	17.72	12	1.2	2.65
	ORF SEQ ID NO:		11782	11811		12389	12479		13902	13911	14709		10516		11418	11466	11932	11933			12187				13572	13966	14016	14208	14394	14481
	SEQ ID NO:		6687	6734		7281		7834	8904	8918	9724	9854	7721	6514	6367	8408	6845	6845		7020	7073	7373	7444	8035	8565	8981	9027	9228	9407	9502
	Probe SEQ ID NO:	1582	1691	1739	2077	2306	2386	2916	3904	3918	4739	4875	487	477	1370	1410	1856	1856		2037	2082	2402	2476	3018	3558	3983	4031	4231	4417	4512

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Single Exon Probes Expressed in HBL100 Cells	Top Hit Descriptor				П	P.chrysosparium lignin peroddase genes, complete cds	gene encoding mitochondrial protein, mRNA	Homo seplens ATP synthase, H+ transporting method.	gene encoding mitochondrial protein, mRNA, mRNA	Starfish (P.ochraceus) cytoplasmic actin gene, complete cds	Mus musculus (CRSwiss chromoldehide a - L	Ureaplasma urealyticum section 57 of 60 of the	ye11g07.r1 Strategens lung (#537210) Home septems CONA classifications	17488 6	Home sapiens hyperpolarization softwated cyclic nucleotide gather profession	PMP-C10400-310700-005-d08 CT0400 Homo sapiens CDNA	FM4-C10400-310700-005-d08 CT0400 Homo series CDNA	Aquifex aeolicus section 7 of 109 of the camplets persons	B.taurus mRNA for D-aspartate oxidase	EST385464 MAGE resequences, MAGM Homo seniens - PAIA	Danio rerio peptide YY precursor gene, complete cole	Arabidopsis thaliana DNA chromosome 4, continuant No. 20	Wg11c07x1 Soares_NSF_F8_9W_OT PA P S1 Home seales_PAYs	Wg11c07.x1 Soares_NSF_F8_BW_OT_PA_P_S1 Homo sapiens cDNA_companiens cDNA_compa	A-AGGLUTININ ATTACHMENT SUBUNIT PRECIDENCE	RHIB PROTEIN	MOLT-INHIBITING HORMONE PRECURSOR MIH)	Choristoneura fumiferana dianausa asserciated	Vibrio cholerae chromosome II, section 73 of 93 of the complete chromosome	Mus musculus annedn V gene, Intron 4 segment containing 5' LTR and den oorlien of N. Edv.	Rathe months. 1777	on70d04.st Scares NEI T 252 2.1	CONTRACTOR OF THE STATE OF THE
Exon Probe	Top Hit Database Source		뒫	EST_HUMAN	EST HUMAN	L.	Z			Z	M	Ę	EST HUMAN	_ <u>t</u>	TOT LINEARY	EST LINAN	NAMOL . C.	ž	Z	EST HUMAN	I N	- 1	- 1	_1.	-1	. 1	ISSPROT					HUMAN	7
Single	Top Hit Acession No.		2 0E-01 AF142703.1	H04858.1	AA884625.1	T.101.5M	4502296 NT	450000	4502296 NT		7	AE002156.1		ARREAGE	BE696604 1	T	T	Ţ.		7	T	T	T	T	Ī				E004416.1	230113.1 NT	Γ		
	Most Similar (Top) Hit BLAST E Value	1 20	7000	2.00.01	2 65 04	N-A-	2.55-01	2.5F-01	2.5E-01			2.5E-01		2.6E-01	2.6E-01	2.5E-01 B	2.5F-01 A	2.5F-01 X		2.5E-01 A		-					2.5F-01 AF	C / <	τι	2.5E-01 AJ	2.5E-01 UB	24E-01 AA936316.1	
	Expression Signal	1 40	37.8	4 83	124		1.57	23	8.7	8	200	14.95		3.61	1.29	1.20	6.55	-	3.33	1.12	6.31	1.55	1.55	1.09	0.91	1.31	4.67	224		3.61	0.8	1.12	
	ORF SEQ ID NO:	14553		ľ	15005	1994	3	10311		10878	-	11142			11922	11823		12646	_	13492	13500	13759	13760				14583	14614	-		14684	10585	
	Exan SEQ ID NO:	9585	L	9826	10038	5301		<b>83</b>	6313	5842	6053	6111		888	1121	/6//	6316	7528	8338	8465	8481	8760	8760	8953	9184	9594	9597	9622	-	200	מנמנ	2000	
	Probe SEQ ID NO:	4577	4812	4877	6909	238	1	2 9	253	822	1043	100	900	3	¥ ;	2 5 6	747	5265	3328	3457	3473	3757	3757	3955	4191	4606	4611	4637	4950	213	1 0 0	5	

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Table 4
Single Exon Probes Expressed in HBL100 Cells

Oryza longistaminata receptor kinase-like protein, family member D, and retrofit (gag/pol) genes, complete od Aquifex secilicus section 12 of 109 of the complete genome 7h23d04.x1 NCI\_CGAP\_Co16 Homo sepiens cDNA clone IMAGE:3316807 3' similar to SW:PRSB\_XENI\_A Marinilabilia agarovorans gyrB gene for DNA gyrase subunit B, partial ods, strain:IFO 14957 no16408.s1 NCI\_CGAP\_Phe1 Homo sapiens cDNA cione IMAGE:1100843 3' similar to contains Alu Mesembryanthemum crystallinum pulative potassium channel protein Mkt1p mRNA, complete cds и97h10.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:213283 6 Oncorhynchus mykiss shaker-related potassium channel Tshaz gene, complete cds #21b07.s1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE:130357 3 1/17/01.r1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE:149017 5 Methenococcus jannaschii section 138 of 150 of the complete genome 601142073F1 NIH\_MGC\_14 Homo sapiens cDNA clone IMAGE:3505818 67 601175562F1 NIH\_MGC\_17 Homo sapiens cDNA clone IMAGE:3531015 5 802132442F1 NIH\_MGC\_81 Homo sapiens cDNA clone IMAGE:4271578 5 Rattus norvegicus mRNA for alphaB crystallin-related protein, complete cds Zaccys dhumnadas fructose-1,6-bisphosphatase mRNA, complete cds IMMUNOGLOBULIN A1 PROTEASE PRECURSOR (1GA1 PROTEASE) Escherichia coli K-12 MG1855 section 202 of 400 of the complete genome Homo sapiens KIAA0851 gene (partial), XT3 gene and LZTFL1 gene Homo sepiens KIAA0851 gene (partial), XT3 gene and LZTFL1 gene aromatase [Poephila guttata≕zebra finches, ovary, mRNA, 3188 nt] Aycoplasma genitalium section 35 of 51 of the complete genome **Top Hit Descriptar** Homo sapiens partial intron 3 of the wild type AF-4/FEL gene epetitive element.contains element THR repetitive element ; 042588 26S PROTEASE REGULATORY SUBUNIT 6A; forno sapiens KIAA0450 gene product (KIAA0450), mRN -ycopersicon esculentum PRF (Prf) gene, complete cds H.sapiens AGT gene, Pstl fragment of Intron 4 Human erythropoletin gene, complete cds Mus musculus cdh5 gene, exon 1, partial Bovine adenovirus 3 complete genome D.discoldeum (Ax3-K) ponA gene S.pombe swiß gene EST\_HUMAN EST\_HUMAN **EST HUMAN** EST HUMAN EST\_HUMAN SWISSPROT Top Hit Detabase È 눋 7662133 Top Hit Acession 2.4E-01 BF002171.1 AA601379.1 24E-01 AE000312.1 2.4E-01 AJ289880.1 AF251708.1 AE000680.1 AF252302.1 2.4E-01 AF267753.1 2.4E-01 AF251708.1 ġ AJ289880. 2.3E-01 R21732.1 2.3E-01 U65391.1 2.3E-01 H69836.1 2.3E-01 Y10887.2 2.3E-01 M11319.1 2.4E-01 X71783.1 2.4E-01 U72726.1 2.3E-01 U67596.1 2.4E-01 P45384 23E-01 2.3E-01 24E-01 2.4E-01 2.4E-01 2.4E-01 24E-01 2.3E-01 2.4E-01 2.4E-01 2.4E-01 2.3E-01 2.3E-01 (Top) HII BLAST E 6.04 0.83 27.33 2.09 1.08 0.91 1.23 1.58 0.92 1.97 6.25 131.89 1.53 0.71 1.38 14.3 14.3 14.57 8 40. 3.8 Expression Signal 3.01 10693 10968 12470 12833 13326 14937 11650 12656 12415 13105 13252 11936 12301 12553 12761 12783 13683 10441 11411 ORF SEQ ÖNQ 7912 8027 8239 7646 7542 6362 6848 5659 5686 6588 6978 7350 SEQ ID 5857 6284 7295 7436 8680 0966 5426 8834 6284 284 8076 5837 8231 6804 8 921 88 2 1592 1994 2893 3010 3216 3288 3832 4224 2467 2688 3675 2378 Probe SEQ ID 1814 2104 3059 3075 1285 1859 2200 2713 3920 927

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Mus musculus renin (Ren-1c) gene, promoter region	Synechocystis sp. PCC8803 complete genome, 1/27, 1-133859	Homo saplens mitogen-activated protein kinase p38detta (PRKM13) mRNA, complete cds	Homo saplens nuclear transport factor 2 (placental protein 15) (PP15) mRNA	Mus musculus tulip 1 mRNA, complete cds	oz14a10.x1 Soares, fetal. liver, spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:1675290 3' similar to TR:013040 Q13040 ATP-BINDING CASSETTE PROTEIN;	Homo sapiens PPAR detta gene, promoter region	Trimeresurus malabaricus cytb gene, partial cds; mitochondrial gene for mitochondrial product	Fresh-water sponge Emf1 alpha collegen (COLF1) gene	602085608F1 NIH_MGC_83 Hamo sapiens cDNA clone IMAGE:4248969 5"	601462629F1 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3866190 67	601462629F1 NIH_MGC_67 Homo sapiens cDNA done IMAGE:3868190 51	PM2-HT0353-281299-003-e12 HT0353 Homo septens cDNA	PM2-HT0353-281289-003-e12 HT0353 Homo sapiens cDNA	Homo sepiens FRA3B common fragile region, diadenosine triphosphate hydrolasse (FHIT) gene, exon 5	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 62	Homo sapiens chromosome 21 segment HS21C085	Xiphophorus maculatus truncated Rext retrotransposon reverse transcriptase (RT) pseudogene	Drosophila melanogaster UNC-119 (unc-119) gene, complete cds	Mus musculus mixed lineage kinase 3 (Mik3) and two pore domain K+ channel subunit (Kcnk5) genes, complete cds	Mus musculus MAP kinase kinase kinase 1 (Mekkt) mRNA, complete cds	Mus musculus MAP kinase kinase kinase 1 (Mekk1) mRNA, complete cds	Human scRNA (BC200 beta) pseudogene	Human scRNA (BC200 beta) pseudogene	Human beta-cytoplasmic actin (ACTBP9) pseudogene	zq87c05.r1 Stratagene hNT neuron (#937233) Homo sapiens cDNA clone IMAGE:648968 5	Mus musculus vinculin gene, exxn 3	yr42h09.r1 Soares fetal liver spleen 1NFLS Home saplens cDNA clone IMAGE:208001 6' similar to gb:214116_ma1 cD59 GLYCOPROTEIN PRECURSOR (HUMAN);	nm31e11.s1 NCI_CGAP_Lip2 Hamo saplens aDNA clone IMAGE:1061804	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 16
Top Hit Database Source	F	N.	<u>F</u>	5	L.	EST_HUMAN	LN	Ŋ	F	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	<b>—</b> ₩	N N	F.	Į.	ΤŃ	<b>_₽</b>	노	N	IN	Z.	NT	EST_HUMAN	TN	EST_HUMAN	EST_HUMAN	LN.
Top Hit Acession No.	.78789.1	D80899.1	AF092535.1	5031984 NT	\B032400.1	N052190.1		AF171901.1	M34640.1	BF677538.1	3E618258.1	SE618258.1		BE155625.1	AF020503.1	AL161562.2	AL163285.2	AF165728.1	AF119102.1	AF166142.1		Γ	U01307.1	U01307.1	D50604.1	AA211218.1	L13299.1	H60548.1	AA569289.1	AL161504.2
Most Similar (Top) Hit BLAST E Value	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.3E-01	2.2E-01		22E-01			2.2E-01	2.2E-01	ľ	2.2E-01	2.2E-01	_	2.2E-01		2.2E-01	2.2E-01		_		2.2E-01			2.2E-01	2.2E-01		
Expression Signal	1.85	0.87	2.08	. 5.79	69.0	0.8	2.42	1.16	1.86	4.2	1.38	1.38	5.48	5.48	1.44	23	1.51	1.61	1.09	6.31	224	224	124	1.24	1.4	2.15	1.24	0.91	1.5	1.79
ORF SEQ ID NO:		14305	14330	14400	14832	10178			12128	12434	12595	12596	12848	12849		•	13676			14078			14206			14651		14915		
SEQ ID NO:	9270	8320	8351	9412	9861	5165	6530	6954	7018	7313	7480	7480	7832	7832	7870	8319	8672	8728	9085	8092	9136	9136	9224	9224	9664	6996	9857	9838	5972	5974
Probe SEQ ID NO:	4277	4329	4360	4422	4882	88	1532	1969	2035	2339	2512	2512	2812	2812	2850	3308	3667	3724	4091	4098	4141	4141	4230	4230	4679	4684	4878	4961	958	928

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Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Chiamydia muridarum, section 45 of 85 of the complete genome	Mus musculus interferon (alpha and beta) receptor 2 (lifnar2), mRNA	Muse muscaling interferon (alpha and beta) receiptor 2 ((fnar2), mRNA	MANUEL HANDEN CONTINUE HANDEN CONTINUE HANDEN CONTINUE TO CONTINUE	ok/3e02.s1 NCI_CGAP_GC4 Fonto 8eptens curva ciglis introduction of community community of the complement cs Precursor (Human);	602083129F1 NIH_MGC_81 Homo septens cDNA clone IMAGE:42475US 5	Homo saplens potassium voltage-gated channel, subfamily H (egg-related), member 4 (KCNH4), mRNA	Beta vulgaris mitochondrion, complete genome	INMEDIATE-EARLY PROTEIN IE180	IMMEDIATE-EARLY PROTEIN IE180	Orchestia cavimana calcium-binding protein BP23 predursor (BP23) gone, compress cus	Homo sapiens mRNA for KIAA1215 protein, partial cds	Homo sapiens pshsp47 gene, complete cds	Homo saplens hax11 proto-oncogene, exans 1 to 3 and hug-1 gene	GENOME POLYPROTEIN [CONTAINS: CAPSID PROTEIN C (CORE PROTEIN) (P22); ENVELOPE CONTAINS (COPS) (M21); PROTEIN	PT: NONSTRUCTURAL PROTEIN NS2 (P21); PROTEASEMELICASE NS3 (P70); NONSTRUCTURAL	PROTEIN>	Human surfactant protein-C (SP-C) gene, complete cds	Gallus gallus mRNA for avena, complete cds	Homo saplens CGI-18 protein (LOC51008), mRNA	O.cunniculus germline IdH heavy chain v-ri pseudogane, anayyo v naz	Mus musculus Major Histocompatibility Locus class II regimi	Synechocystis sp. P.C.Cosus complete genome, 1121, 101745 sectors	Homo sapiens chromosome 21 seginein no.4 100 13	Homo sapiens fact gene	PM1-H10422-281285-02-00 F10424 Tulib aquale way	Plum pox wrus strain m, continue general, social continue, social continue and social	Home sapiens dysucations, aimia (2 may, minor)	Home sapiens mixed, on this same and a partial cols	Home septems scalutivates symptom mirror, per an ore	Human Disaykinii bi Tecepini (staajbr) Bain, compress com	Homo saplers 14432 dagged years, conpress cost and
	Top Hit Detsibase Source	F				EST_HUMAN	EST HUMAN	— <u>E</u>	П	SWISSPROT	SWISSPROT	IN	NT	N	NT			SWISSPROT	NT	NT	NT	L	FZ	LN L	Į.	Į,	EST HUMAN	Z.	ŁZ	Į.	Ę	Ę	<u> </u>
	op Hit Acession No.	E002314.2	TA 200 NT	1 0004-000	1 N 8876/9	2.1E-01 AA906824.1 E	П	6912445 NT	1N 19838361 NT	911676	<b>211675</b>	1F124528.1	\B033041.1	AB010273.1	1,1009794.1		,	528880	<u>-</u>	AB017437.1	TV05601 NT	M77085.1	AF027865.1		AL163213.2	AJ132695.5	AW384937.1	AJ24395	4503408 NT	AB007974.1	AF280700.1	U22346.1	AF111170.3
-	Most Similar (Top) Hit BLAST E	2.1E-01 A	200	2.15-01	2.1E-01	2.1E-01	2.1E-01	2.1E-01	2.1E-01	2.1E-01	2.1E-01	2.1E-01 /	2.1E-01	2.1E-01	2.1E-01			2.15-01	2.1E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01				2.0E-01			2.0E-01
	Expression Signal	2.16		72	1.22	1.62	242	2.25	8.22	1.28	1.28	0.9	1.28	1.86	4			0 0	F	284				0.68	2.91	1.34		1.01	7.19	3.81	1.54	1.48	1.48
	ORF SEQ ID NO:	1		11217	11218	11946		12889		13930				14416					15003			10728			11144				11612	11574	L		
	SEQ ID	R113	1		6181	8858		7874	L		L	L		L				7000	L	4		L	L	L					L	L		L	Ш
	Probe SEQ ID NO:	4407	5	=	1179	1860	2094	2854	3718	3940	3940	4128	4251	4443	4719			-	5087		230	S C	\$ 2	8	1108	1235	1289	1432	1456	1520	1526	1658	1679

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Methenococcus Jannaschil section 67 of 150 of the complete persons	Homo sabiens hypothetical protein F. 110120 (F) 1101201 mRNA	H.saplens Na+D-ducose cotransport required game	HOMEOBOX PROTEIN GLABRA2 (HOMEOBOX-LEUCINE ZIPPER PROTEIN ATHB-10) (HD-ZIP PROTEIN ATHB-10)	xp15b02.x1 NCI_CGAP_HN9 Homo septens cDNA clone IMAGE:2740395 3' strailar to contains element. MER21 resettitive element:	CED-11 PROTEIN	C.parasitica eanC rene	QV4-EN0032-190500-223-e03 EN0032 Homo sepiens cDNA	Homo sapiens hypothetical protein ASH1 (ASH1), mRNA	Homo saplens putative psthHbD pseudodene for hair keratin, excess 1 to 9	Homo sepiens full length Insert cDNA YH85A11	Mus musculus cubilin mRNA, partial cds	yb17a10.r1 Stratagene fetal spleen (#937205) Horno sablens cDNA clone IMAGE:71418 5	Rattus novegicus Aryl hydrocarbon receptor nuclear translocator 1 (Amrt1), mRNA	Mus musculus pale ear (ep) gene, wild type allele, 3' region, partial cds	Homo saplens lambdailota protein kinase C-interacting protein mRNA, complete cds	Homo saplens lambda/lota protein kinase C-interacting protein mRNA, complete cds	RC3-BT0502-251189-011-401 BT0502 Homo sapiens cDNA	RC3-BT0502-251189-011-d01 BT0502 Hamo sepiens cDNA	Mus musculus interteukin 2 receptor, gamma chain (II2rg), mRNA	EST67784 Fetal tung II Homo sapiens cDNA 5' end	Sorghum bicolor 22 kDa kafirin cluster	Plasmodium wwax reticulocyte binding protein-2 (rbp-2) gene, complete cds	Homo saplens hypothetical protein FLJ10581 (FLJ10581), mRNA	Sigmodon hispidus p53 gene, partial cds	Gallus gallus ovalbumin (Y) gene, complete cds	Mouse gene for immunoglobulin diversity region D1	y42/10.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:129547 6	Rattus norvegicus arylacetamide deacetylase gene, complete cds	PAIR-RULE PROTEIN ODD-PAIRED	Schizosaccharomyces pombe DNA for cytoplasmic dynein heavy chain, complete ods	CM3-CT0315-271199-045-b11 CT0315 Hamo saplens cDNA
Top Hit Datebase Source	¥	¥	N.	SWISSPROT	EST HUMAN	SWISSPROT	LZ	EST HUMAN	NI	Ę	FZ.	Ę	EST_HUMAN	LN T	NT	NT	NT	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	NT	NT	NT	NT	NT	NT	EST_HUMAN	NT	SWISSPROT	L	EST_HUMAN
Top Hit Acession No.	J67525.1	8922238 NT	<b>G2877.1</b>	246607	AW238005.1	234641	(83997.1	3E826165.1	8922080 NIT	/19216.1	2.0E-01 AF074990.1	\F197159.1		7549743 NT	\F004363.1	J32581.2	J32581.2 ·	3E070801.1	E07080	7305180 NT		F061282.1	\F184623.1	8922533 NT	J66066.1	100922.1	13197.1	116467.1	1.9E-01 AF264017.1			1.9E-01 AW754108.1
Most Similer (Top) Hit BLAST E Vefue	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01	2.0E-01 A	2.0E-01	1.9E-01	1.9E-01	1.9E-01 U	1.9E-01	1.9E-01	1.9E-01	1.9E-01	1.9E-01	1.9E-01 /	1.9E-01 A	1.9E-01	1.9E-01	1.9E-01	1.9E-01 D13197.1	1.9E-01 R16467.1	1.9E-01	1.9E-01 P39768	1.9E-01 /	1.9E-01/
Expression Signal	3.96	0.93	1.21	. 0.68	0.67	0.7	0.77	9.72	7.34	0.93	7.46	1.19	1.29	9.3	5.34	24.68	24.68	6.04	5.32	0.96	. 12.01	3.22	2.64	2.22	3.91	5.5	4.28	4.85	0.77	-	3.37	1.47
ORF SEQ ID NO:		11930		13444		13637	13912			14885		14985			ĺ			10690	10690			11400		12414	12887		13345	13434	13721	13749	13886	13956
Ean SEQ ID NO:			7262	8416	8497					8086	7840	10016	10029	5182	5402	9299	9299	5683	5683	5986	6094	6350	6407	7294	7872	7887	8322	8408	8720	8750	8887	8967
Probe SEQ ID NO:	1720	1853	2286	3407	3489	3624	3919	4433	4874	4931	4979	5045	5059	110	320	648	648	655	656	971	1087	1353	1409	2319	2852	2868	3311	3388	3716	3746	3886	3968

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	MR1-FN0010-290700-007-404 FN0010 Home seralens cDNA	Rattus novegicus chemokine receptor CXCR3 mRNA, complete cds	Mus musculus p116Rip mRNA, complete cds	Mus musculus Cotg gene for chaperonin containing TCP-1 gamma subunit, partial cds.	Homo sapiens calcium channel, voltage-dependent, beta 2 subunit (CACNB2) mRNA, and translated	products	Original rappes grate for membrane guantyly cyclese Origin, complete cds wd71f02 vt NCI CGAD 1.124 Home contene of NA Alma 14A CE-2232724 4	Dichostalium discodeum plasmid Ddo5, complete cenome	Yersinia pestis plasmid pCD1	Homo sapiens latent transforming growth factor beta binding protein 4 (LTBP4) mRNA	9922410.55 NCI_CGAP_Kid3 Home septens cDNA clone IMAGE:1761611 3' similar to TR:075936 075636 GAMMA BUTYROBETAINE HYDROXYLASE:	Mus musculus Scyed, Scye16-ps, Scye5 genes for small Inducible cytoldre A6 precursor, small	Inducible cylokine A9 precursor, Scya16 pseudogene, email inducible cylokine A5 precursor, complete cds	QV3-DT0018-081289-036-g04 DT0018 Home septems cDNA	Jonopskijum acaule LEAFY protein (LEAFY2) gene, partial cds	x41a03.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2659766 3'	QV0-BN0041-070300-147-c04 BN0041 Hamo sapiens cDNA	601809723R1 NIH_MGC_18 Hamo sepiens cDNA dane IMAGE:4040621 3'	1/45e01.s1 Soares placenta Nb2HP Homo saplens cDNA clone IMAGE:151704 3' similar to contains Alu repetitive element:	1/45601.s1 Soares placenta Nb2HP Homo septens cDNA clone IMAGE:1617043' similar to contains Alu	repetitive element;	Bovine NB25 mRNA for MHC class II (BoLA-DQB), complete cds	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 56	Mus musculus Scyas, Scyar8-ps, Scyas genes for small inducible cytokine A8 precursor, small	indúcible cytokine Av precursor, Scyarl 6 pseudogene, small inducible cytokine A6 precursor, complete cds	S.tuberosum mRNA for alcohol dehydrogenase	MR3-ST0203-151289-112-g06 ST0203 Homo saplens cDNA	Mesocricetus auratus Na-taurocholate cotransporting polypeptide mRNA, partial ods	ti57e04.x1 NCI_CGAP_Lym12 Homo saplens cDNA clone IMAGE:2134590 3'
Top Hit Detabase Source	EST HUMAN	뉟	본	Į.		Z	EST HIMAN	L	NT	NT	EST_HUMAN		NT	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN		EST_HUMAN	NT	NT	!	LN	NT	EST_HUMAN	NT	EST_HUMAN
Top Hit Acession No.	3E834943.1	AF223642.1	J73200.1	AB022080.1		1 N 2882082 IN 1	19122121	AF000580.1	AL117189.1	4505036 NT	1733708.1		VB051897.1	W835728.1	\F184589.1	W182300.1	W995178.1	3F183582.1	H03369.1		103369.1	37954.1	\L161558.2		-	02179.1	W814270.1	\F181258.1	1439881.1
Most Similar (Top) Hit BLAST E Value	1.86-01	1.9E-01	1.8E-01	1.8E-01	70 20 7	1.85-01	1.8E-01	1.8E-01	1.8E-01	1.8E-01	1.8E-01		1.8E-01	1.8E-01	1.8E-01	1.8E-01	1.8E-01	1.8E-01	1.8E-01		1.85-01	1.8E-01	1.8E-01		1.8E-01 /	1.8E-01	1.8E-01	1.8E-01 /	1.8E-01
Expression Signal	-	1.11	1.73	. 1.97	97.7	8 5	0.75	1.17	7.26	1.18	1.34		1.42	1.63	1.6	1.28	2.5	99.0	0.78		0.78	1.21	6.34		3.53	1.89	2.79	6.38	1.04
ORF SEQ ID NO:	14090			10327	40.493				11309				11948					13322	13553		13554		14401	,	14601	14634	14830	14877	14899
Exan SEQ ID NO:	9104	9833	5112	7717	8448	5760	5982	8082	6267	6800	6819		6860	7582	7848	7853	8068	8296	8546	1	200	9 8 8	9414		9612	9646	9859	9901	8919
Probe SEQ ID NO:	4110	4851	32	258	380	737	296	1075	1269	1810	1829		1871	2620	2827	2833	3051	3285	3540	1	200	4206	4424		4	4681	4880	4924	4942

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Homo saplens derivative 11 breakpoint fragment: partial intron 10 of the ALL-1/MIL/HRX gene fused to intron qn57e09.x1 Soares\_fetal\_liver\_spleen\_1NFLS\_S1 Homo saplens dDNA done IMAGE:1848808 3' sImiliar to Taxus canadensis gerany/gerany/ diphosphate synthase mRNA, complete cds Anabeena sp. ORF4 (partial), ORF3, ORF3, ORF7, adpA gene, adpB gene, adpC gene, adpD gene, adpE Vibrio cholerae hypoxanttine phosphoribosyltransferase (hpt) gene, partial ods, hemegglutinin/protease regulatory protein (hapR) gene, complete ods, and YRAL VIBCO gene, partial ods EST41651 Endometrial tumor Homo saplens cDINA 6' end Vibrio choleras hypoxanitine phosphoribosytransferase (hpt) gene, partial cds, hemegglutinin/protesse regulatory protein (hspR) gene, complete cds, and YRAL VIBCO gene, partial cds P.dumariii histone gene cluster for core histones H2A, H2B, H3 and H4 NEUROFILAMENT TRIPLET I PROTEIN (NEUROFILAMENT LIGHT POLYPEPTIDE) (NF-L) Lymantria dispar nucleopolyhedrovirus, complete genome Arabidopsis thaliana DNA chromosome 4, config fragment No. 69 Homo sepiens BNIP3H (BNIP3H) gene, complete cds; nuclear gene for mitochondrial product yh76f12.r1 Soares placenta Nb2HP Homo sepiens cDNA clone IMAGE:135699 5' nk28d12.s1 NCI\_CGAP\_Co11 Homo sepiens cDNA clone IMAGE:1014839 3' 601274604F1 NIH\_MGC\_20 Homo septems oDNA clone IMAGE:3615768 6 Homo sapiens homeobox protein OTX2 gene, complete cds. AXONIN-1 PRECURSOR (AXONAL GLYCOPROTEIN TAG-1) Crassostrea gigas RNA polymerase II largest subunit mRNA, partial cds. **Top Hit Descriptor** Lymantria dispar nucleopolyhedrovirus, complete genome contains OFR.b1 OFR repetitive element; Homo sepiens mRNA for KIAA0472 protein, partial ods Homo sapiens mevalonate kinase gene, exon 6 and 7 Escherichia coli reverse transcriptase, retron EC86 Escherichia coli reverse transcriptase, retron EC86 Schistocerca gregaria alpha repetitive DNA Homo sapiens hap1 gene, complete CDS Naja naja atra ctx-1 gene, exons 1-3 Naja naja atra ctx-1 gene, exons 1-3 5 of the AF-4/FEL gene gene and adpF gene NT SWISSPROT EST HUMAN EST HUMAN **EST HUMAN** EST\_HUMAN EST\_HUMAN SWISSPROT Top Hit Database Saurce 닐 뉟 Ę 눋 Ę 뉟 Þ Top Hit Acession 1.7E-01 AB007941.1 1.6E-01 AF217532.1 AF298117.1 AA548863.1 AL161573.2 1.7E-01 AJ238736.1 1.7E-01 AF081514.1 1.7E-01 AF081810.1 1.7E-01 AF081810.1 1.7E-01 AF255051.1 1.7E-01 AF000718.1 1.7E-01 AF000716.1 1.7E-01 AA336909.1 1.7E-01 AJ238736.1 AJ269505.1 AJ224877.1 AJ235377. 1.7E-01 AI247635.1 1.7E-01 BE385164.1 1.7E-01 X53330.1 호 1.6E-01 R31497.1 1.7E-01 X52936.1 U10334.1 1.6E-01 P22063 1.8E-01 X60208. 1.7E-01 P35616 1.6E-01 1.7E-01 1.6E-01 1.7E-01 1.7E-01 (Top) Hit BLAST E Value Aost Simila 1.31 1.58 1.11 0.6 8 8 1.45 6.17 1.89 0.93 2.05 1:17 1,92 1.31 1.31 1,96 Expression 11542 10208 10705 11529 12965 13058 13540 13843 14659 11961 1086 88 12826 12827 12892 12964 13391 10848 14998 10603 11081 11865 14997 ORF SEQ ÖNQ 9676 5194 7695 6872 6924 7947 8049 8533 6051 7809 7809 7876 9418 6487 10030 8371 8837 SEQ ID 10030 5605 5817 6051 6921 3835 4932 126 671 1474 1490 1883 1938 3363 5061 2788 SEQ ID 1946 1946 1946 1935 2788 2928 3527 4428 1787 1781 2828 3032 ÿ

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	H. sapiens mRNA for novel T-cell activation protein	Homo sapiens mRNA for KIAA1308 protein, pertial cds	Homo sapiens cytochrome P450 3A4 (CYP3A4) gene, promoter region	Homo sapiens cytochrome P450 3A4 (CYP3A4) gene, promoter region	Populus trichocerpa cv. Trichobel ABI3 gene	Populus trichocarpa cv. Trichobel ABI3 gene	Archaeoglobus fulgidus section 145 of 172 of the complete genome	Vibrio cholerae chromosome II, section 70 of 83 of the complete chromosome	Homo saplens apelin gene, complete cds	EST380877 MAGE resequences, MAGJ Hamo sapiens cDNA	Mus musculus cheperonin subunit 3 (gamma) (Cct3), mRNA	284h09.s1 Strategene colon (#937204) Homo saplens cDNA clone IMAGE:511361 3' similar to TR:E221955 E221955 39,855 BP SEGMENT OF CHROMOSOME XIV.;	Lycopersicon esculentum Rsal fragment 2, satellite region	Lycopersicon esculentum Real fragment 2, satellite region	LOW-DENSITY LIPOPROTEIN RECEPTOR-RELATED PROTEIN 2 PRECURSOR (MEGALIN) (GLYCOPROTEIN 330)	IL3-HT0619-040700-197-E05 HT0819 Homo sepiens cDNA	IL3-HT0619-040700-197-E05 HT0619 Homo saplens cDNA	AV711698 DCA Hamo seplens cDNA clane DCAADH06 5'	Homo sapiens chromosome 21 segment HS21C084	Homo saplens partial SLC22A2 gene for organic cation transporter (OCT2), exon 1	Rattus norvegicus Insulin-responsive glucose transporter (GLUT4) gene, 5' end	xn39d11.x1 NCI_CGAP_Kd11 Homo sapiens cONA clone IMAGE:2696085 3*	Human gene for dihydrolipoamide succlnyfransferase, complete cds (exon 1-15)	Human gene for dihydrolipoamide succinyfransferase, complete ods (exon 1-15)	Mus musculus MAP kinase kinase kinase 1 (Mekk1) mRNA, complete cds	602083269F1 NIH_MGC_81 Homo saplens cDNA clone IMAGE:4247537 67	xw58a02.x2 NCI_CGAP_Pan1 Homo saplens cDNA clone IMAGE:2831978 3' similiar to gb:X55072_ms1 THYROID HORMONE RECEPTOR ALPHA-1 (HUMAN);	oo88d05.s1 NCI_CGAP_GC4 Homo sepiens cDN4 clone IMAGE:1671337 3' similar to gb:M11433 RETINOL-BINDING PROTEIN I, CELLULAR (HUMAN);	Mus musculus ICR/Swiss glyceraldetyde 3-phosphate dehydrogenase (Gapd-S) gene, complete ods
Top Hit Database Source										EST HUMAN		EST_HUMAN			SWISSPROT	EST_HUMAN	EST_HUMAN	EST_HUMAN				EST HUMAN				EST HUMAN	EST_HUMAN	EST_HUMAN	-
	支	Ę	Þ	Ĭ	¥	Ę	ΙN	ĹΝ	N	EST	ĽΝ	EST	Ę	ΝŢ	MS	EST	EST	EST	Ż	F	Ż	13	Ę	Ż	¥	EST	EST	EST	눌
Top Hit Acession No.	<b>04232,1</b>	AB037729.1	1.6E-01 AF185589.1	\F185589.1	1.6E-01 AJ003165.1	1.6E-01 AJ003165.1	1.6E-01 AE000982.1	1.6E-01 AE004413.1	1.6E-01 AF179680.1	AW968601.1	6753319 NT	AA088343.1	AJ006358.1	1.6E-01 AJ006356.1	298158	1.5E-01 BE710087.1	1.5E-01 BE710087.1	1.6E-01 AV711696.1	1.5E-01 AL163284.2	AJ251885.1	1.6E-01 L38125.1	1.6E-01 AW 195516.1	1.5E-01 D26535.1	1.6E-01 D26535.1	AF117340.1	1.5E-01 BF695381.1	1.5E-01 AW 572518.1	AA835049.1	U08964.1
Most Similer (Top) Hit BLAST E Value	1.6E-01 XB	1.6E-01 AE	1.6E-01	1.6E-01	1.6E-01	1.6E-01	1.6E-01	1.65-01	1.6E-01	1.8E-01	1.6E-01	1.6E-01	1.6E-01		1.6E-01 P	1.5E-01	1.5E-01	1.6E-01		1.5E-01								1.5E-01	1.6E-01
Expression Signal	0.98	1.22	76.01	10.97	1.18	1.16	0.73	2.51	10.56	2.61	3.9	1.47	1.19	1.19	121	1.38	1.38	2.11	1,39	1.81	2.69	0.7	2.85	2.85	1.47	1.34	1.06	4.64	
ORF SEQ ID NO:	12418	12514	12859	12860	13562		13692		14177			14765	14776	14777	14973			L	10824	11116		11234	11295	11296	11509	12713		13310	
Exan SEQ ID NO:		7383	7842	7842				8894	9195	8325	9331	9772	9794		10000	1		L	5786	6087		6198	6254	6254	8448	7599	7863	8286	
Probe SEQ ID NO:	2322	2422	2821	2821	3548	3548	3685	3894	4202	4334	4340	4788	4810	4810	5020	246	246	583	774	1080	1096	1197	1256	1256	1451	2639	2843	3274	3670

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Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo sepiens pyruvate dehydrogenase kinase, iscenzyms 1 (PDK1), nuclear gene encoding mitochondrial metals.	DN es	RC2-HT0149-191000-043	Homo seniens chromosome 21 secretari USA Chou	602087192F1 NIH MAC 67 Home emine and a line in the progression and a line in the line in the progression and a line in the li	602083269F1 NIH MGC 81 Home septients CUMA clothe IMAGE: 4066223 6	Arabidonais thatiana DNA chamanana 4	Homo sacions Total prompts have been a TODA VOSED L. TODA VOSED L.	Xenous leads mRNA for DNA (mens, 1 on by 6337 to 1 or By 2152A2 region	vd54c01 s1 Scares fetal liver snices 1NEI S Lower series - Child Lives -	Mile miseratus around differentiation in the control of the Contro	Themotions maritims seed 22 of 438 of the	NV72407 et NCI CCAB CCP4 Liver Campings gandme	wm/24/01 x1 NCI COAP 112 U	602013527F1 NCI CAAD B 84 U	VOSTA11 Series Infant Ivola ANIB U.	vg/7803 r1 Sharas Infant broin 1NID Linns capiens CDNA	658-02-xf NCI CGAP 1.024 Homo serviers CONA clone INAGE:41487 6	b56c02.xf NCI CGAP 1124 Home enters child Act 114 CE 2010 3	Thermotoga maritima section 22 of 138 of the complete section.	Homo septens G protein-coupled recentor 50 (GPR50) mRNA	Homo saplens G protein-coupled recentr 50 (GPR50) mRNA	Homo saplens gene for NBS1, complete cds	Human calicivirus HUNLV/Girlington/83/UK RNA for canaid protein (CIBES)	Himan calicidate til IAN Worth and Month of the Data A	Polymeriii histore ages chiche to con litter to the transmitted to the transmit histore ages chicken to the transmitted to the	Rettus novembers & kinese enchor models made a constitution of the	Botrytis cheese strain T4 cDNA library inder conditions of alternations don't have	Bothytis gineres strein T4 cDNA library inder conditions of all co	AV712467 DCA Home septens cDNA close DCAAEFG 6	Homo sapiens adapter protein CMS mRNA, complete cds.
	Top Hit Database Source	 	EST HIMAN	EST HIMAN	LN	EST HUMAN	EST HUMAN	Į.	N.	Ė	EST HUMAN	Ę	IN	EST HIMAN	EST HIMAN	EST HIMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	N	II.	H	M	¥	<u> </u>	L	Ę	L	LN	HUMAN	Т
1.6	Top Hit Acession No.	7108358 INT	AW665983.1	Ī	T	Γ		Γ		Ī		5679980	4E001710.1	T	T	F341524.1	59232.1	59232.1	699094.1	1699094.1	E001710.1	4758467 NT	4758467 NT	B013139.1	J277606.1	1277808 1	63330.1	F139518.1				
	Most Similar (Top) Hit BLAST E Value	1.6E-01	1.年点	1.5E-01	1.5E-04	1.6E-01	1.5E-01	1.5E-01	1.4E-01	1.4E-01	1.4E-01	1.46-01	1.4E-01	1.4E-01	1.4E-01	1.4E-01 B	1.4E-01 R	1.4E-01 R	1.4E-01 A	1.4E-01 A	1.4E-01 A	1.3E-01	1.3E-01	1.3E-01 A	1.3E-01 A	1.3E-01 A	1.3E-01 X	1.3E-01 /	1.3E-01 /	1.3E-01	1.3E-01	1.3E-01
	Expression Signal	1.04	2.38	0.96	8.83	1.41	2.83	1.66	1.61	2.72	2.11	1.2	1.61	8.74	4.16	1.65	1.17	1.17	11.16	11.18	3.6	2.28	2.28	1.88	1.51	1.61	0.83	1.49	1.51	2.67	1.71	0.91
	ORF SEQ ID NO:	13691	13782	13929				14799					11783		12701		13811	13812		14035	14094	10384	10385	10563	10663	10864	10893	10942	11049		11233	
	SEQ ID NO:			8936					5354	6917		6704		6925	7877	8472	8806	8806	9046	9046	9109	5375	5375	5560	5658	9858	6853	5901	6020	6116	. 6197	6415
	Probe SEQ ID NO:	3684	3775	3937	4059	4585	4607	4840	297	888	1240	1709	1712	1939	2720	3464	3803	3803	4050	4050	4115	88	g	625	88	880	88	883	1010	1110	1196	1418

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Single Exon Probes Expressed in HBL100 Cells

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Consider and the constant of t	Top Hit Descriptor	PATAGE OF STATE IN THE STATE OF STATE O	Rhodopseudomonas addophila pueB5, pueA6, pueB6, pueB6, pueA6, pueB7, pueA7, pueB8, pueA8 and pueC	POLICIA TO COLOR AND LICENSES.	Archaeolichis fündum million 2 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	Carassius aurahus keratin hara I mDNA	Homo sapiens transcription factor IGHM enhancer 3, JM11 protein, JM4 protein, JM5 protein, T54 protein, JM10 protein, A4 differentiation-dependent protein, triple LIM domain protein 6, and synaptophysin genes.	complete cas, and L-type calcium channel a>	Bowne branched chain alpha-teto acid dihydrolipoyl transacylase mRNA, complete cds	Pyrococcus horikoshii OT3 genomic DNA, 1-287000 nt. position (1/7)	Pyrococcus horikoshii OT3 genomio DNA, 1-287000 nt. position (1/7)	Arabidopsis malana DNA chromosome 4, contig fragment No. 77	munical calicarius HU/NLY/Giffington/83/UK RNA for capsid protein (ORF2), strain HU/NLY/Girlington/83/UK	Human calicivirus HUNLV/Girlington/93/UK RNA for capsid protein (ORF2), strain HUNLV/Girlington/93/UK	becampfilige S/PCZ complete genome	4V3-D10018-081289-036-e03 DT0018 Homo saplens cDNA	Scriistosoma mansoni fructose bisphosphate aldolase mRNA, complete cds	XX23TUXT Scares_NFL_T_GBC_S1 Homo septens cDNA clone IMAGE:2813995 3'	Form equals chronosome 21 segment HS21C080	ob 1 zousor I nirt_McC_9 Home sapers cDNA clone IMAGE:2890063 6	#39602.x1 NCI_CGAP_Brn23 Homo septens cDNA clone IMAGE:2098539 3' similar to gb:U05760_ma1 ANNEXIN V (HI MAN):	Dichostalum discoldeum ORF Do 1018 gene soutel sub-	Home sanjens colon cancer actions and NO As Colon and Colon	AU199146 NT2RM4 Home saniers CDNA class NT2DN44054604 of	AU149148 NT2RM4 Homo septems CDNA class NT2B1400464	AV735249 cdA Homo saplens cDNA clone cdAA_R1167	a48809.st Soares NFL_T_GBC_S1 Hömo sapiens cDNA done IMAGE:14606843' similar to TR:Q16671 Q16871 ANT-MULERIAN HORMONE TYPE II RECEPTOR PRECURSOR	
99001	Top Hit Database Source			EST LIMAN												ESI HUMAN	COT LIMANI	NAMOL	COT DI BAARI	EST HUMAN	EST HUMAN			EST HUMAN	T HUMAN	HUMAN		
•	<u> </u>	Ę		ļű		Ξ	5	#		=		2 2	<u>                                     </u>	z !		3 5	z ů	3 5	i li	3 8	ES.	Ę	ż	S	EST	ES	ES	
	Top Hit Acession No.	AL117078.1	A.1243578 4	1.3E-01 AW812104 1	AE001018.1	1.3E-01 M86918.1	AF108770 4	M24572 4	1.3E-04 APODO004 4	A DOCCOO. 1	Al 484504.0	13F-01 A 1977ans 4		1.3E-01 AJZ/ /606.1	MASSAS 44 4	1.3E-01 AW 304341.1	AW272744 4	AI 183280 2	13E-01 RE2723304	1.3E-01 Al432531.1	1.2E-01 AI421744.1	1.2E-01 U66912.1	NF039442.1	1.2E-01 AU149148.1	NU149146.1	V735249.1	1.2E-01 AA897474.1	
	Most Similar (Top) Hit BLAST E Value	1.3E-01 AL	135-01	1.3E-01	1.3E-01	1.3E-01	135-01	135.01	1 3F-04	4 35 04	4 3E 04	13F-01	100	1 25 01	4 25 04	135.01	13F-01 AW	1.3E-01 AI 1	13F-01	1.3E-01	1.2E-01	1.2E-01	1.2E-01	1.25-01	1.2E-01/	1.2E-01 AV7	1.2E-01	
	Expression Signal	1.66	0.97	1.04	2.79	1.78	86.0	18	0.81	80	137	2	Ę	082	4.10	7 7	1831	1.36	277	1.78	9.01	1.74	2.9	2.5	2.5	3.56	1.03	
	ORF SEQ ID NO:	11999			H	12593	13316	13396	13851	13652		10683	10884			14028	14042		14378	14897	10474	-	-	11405	11406			
	SEQ ID NO:	6905	7086	Li		7478	8290	8376	L	8645	l	8658	5658	9012	9030	8038	9055	9180	9394	9917	5457	5079	5577	6355	6355	6361	6474	
	Probe SEQ ID NO:	1919	2106	2228	2318	2510	3278	3368	3639	3639	3882	3838	3938	4016	4634	4042	4061	4187	4403	4940	382	421	543	1358	1358	1364	1477	

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Table 4
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Single Exon Probes Expressed in HBL100 Cells	Top Hit Descriptor	NUCLEAR FACTOR OF ACTIVATED T-CELLS, CYTOPLASMIC 4 (T CELL TRANSCRIPTION FACTOR NEATS) (NEATS) (NEATS) (NEATS) (NEATS)	A STATE OF S	Heavier INV. Conf. Estat France Begins CDNA Cone IMAGE: 1960553 3'	1. Septemblish for chroogenous retroving like element UHH-Bisakka-10-0-11 of NOT 20-0-0-1-11	AN 1824 567E4 NILL MCC SO U	OV3-BN0046-22030-1-1-0-1-2-4-0 BN0048-U	Human E1A enhancer binding probain (F1A-E) mBNA model = 2.	as80009.x1 Barstead colon HRB7 Home septens CDNA clone INAGE:2335024 3' etnilar to gb:L05095	Human creatine kinese-B mBNA complete cut	Wheet mRNA for a group 2 left of the control of the	OV4-BTOSE Serios As a brase in the property of the property of the A	Mathemasses Increased the Control of	Wheel mRNA for a grown 3 later and 142 of 160 of the complete genome	Wheat mRNA for a group 3 lets emint yegenesis abundant protein (LEA)	Bacillus suhidis complete gasses (need of the fact of	P. Clarkii mRNA recent rection (in 31/077)	P. clarkij mRNA: repeat recion (ID 24/0717)	Rana ridibunda pituitary adenylate cyclase-activating polypeptide variant 2 precursor, mRNA, complete cds, afternatively entradi-	602135185F1 NIH MGC 84 Home series a PMA Ales MACE ASSASSES	Homo saplens chromosome 21 serment HS217n27	Homo saplens chromosome 21 segment HS2/Cn27	RIBONUCIEASE HII (RNASE HII)	h18d08.x1 NCI_CGAP_Bm25 Homo saciens cDNA clave IMAGE:0187089 9	nm08g11.s1 NCI_CGAP_Co10 Homo sepiens cDNA clone IMAGE:1058820 3' similar to gb:X06985_me1 HEME OXYGENASE 1 /LI IMAAN.	60212884751 NIH MGC 8 Home conference (1971)	Arabidoosis thallana DNA chromosome 4 continue formatting and	EST384142 MAGE reseguences, MAGL Home seriens citina	Synechocystis sp. PCC6803 complete genome, 23/27, 28/87/67, annogas	AU140363 PLACE2 Homo seplens cDNA clone PLACE2000403 6	Mus musculus pre T-cell antigen receptor alpha (Ptora), mRNA
XOU PIODES	Top Hit Database Source	SWISSPROT	EST HIMAN	N P	EST HUMAN	EST HIMAN	EST HUMAN	N	EST HIMAN	L	Į.	EST HIMAN	LIN	LN	L	Ę	Z	¥	Ę	EST HUMAN	F	¥	SWISSPROT	EST_HUMAN	EST HUMAN	Т	Т	EST HUMAN	Ę	EST_HUMAN	
aigino	Top Hit Acession No.	014884	2.1	X89211.1	AW449368.1	BF248490.1		Γ	AI720470.1			120	T						AF221633.1	BF677357.1	1.2E-01 AL163227.2 N	163227.2	57599	561003.1	1.1E-01 AA569008.1	Γ				1.1E-01 AU140363.1	6755215 NT
	Most Similar (Top) Hit BLAST E Velue	1.25-01	1.2E-01	12E-01	125-01	12E-01	1.2E-01	1.2E-01	1.2E-01	1.2€-01	1.2E-01	1.2E-01	1.25-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.2E-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01/	1.1E-01 D64004.1	1.1E-01/	1.15.01
	Expression Signal	1.18	2.24	9.87	3.06	231	1.4	1.4	2.31	3.26	6.0	1.63	0.79	0.67	0.67	0.71	1.91	1.91	0.92	9.45	3.98	3.98	4.85	0.74	6.71	1.16	1.48	3.26	1.76	1.79	231
	ORF SEQ ID NO:	11649	11671			12211	12598	12812	12872	12904	12973	13188		13476	13477		14038	14039		14811	14892	14893		10594	10840	11077		11177	11274	11540	
	SEQ ID NO:	6587	6807	6726	П	-	-	7780	7852	7884	7955	8176	8199	8450	8450	8407	8052	8052	9633	8838	8914	8914	10027	5594	5638	6047	2209	146	8228	6485	7227
	Probe SEQ ID NO:	1691	181	1331	1876	2718	2514	2769	2832	2884	2836	3159	3183	3442	3442	3525	4058	4058	4648	4857	4837	4837	2027	88	611	1037	1069	144	<u>8</u>	- -	2250

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Rettus norvegicus Procollagen II alpha 1 (Col2a1), mRNA	Inharten den 12 n3B subunit Imbe. Genomic. 700 nt. segment 4 of 61	I I I I I I I I I I I I I I I I I I I	HSCIRFUZZ ROMBIZBO INIGIN CANA TIGIN SANCES CONTRACTOR	Mus muscutus calcium channet, vortage-dependent, 1 type, apra 15 augum (vectar 19), mon	601308678F1 NIH_MGC_44 Hamo saplens cDNA clone IMAGE:3627066 5	C.reinhardtii nuclear gene on linkage group XIX	yq62g08.s1 Soares fetal liver spleen 1NFLS Hamo saplens cDNA clone IMAGE:200414 3' similar to contains Alu repetitive element;	Alimmersus gene for transposase		G.gailus gene encoding non-histone chromosomal protein HMG-14b, exons 4 and 5	MR3-ST0280-280100-025-907 ST0280 Homo sepiens cDNA	MR3-ST0290-290100-025-407 ST0290 Hamo sapiens cDNA	Drosophila melanogaster klarsicht protein (klar) mRNA, complete cds	Tapa-1≕integral memprane protein TAPA-1 [mice, B cell lymphoma line 38C13, Genomic, 1973 mt, segment 1 - 77	W. I	Aliming the grief of the spicease	WV14h02x1 NCI_CGAP_Birzs nome superis conv. cigne invivor:	Homo sapiens hypothetical protein FLJ20342 (FLJ20342), mrNA	DEOXYRIBONUCLEASE II PRECURSOR (UNASE II) (ACID UNASE) (L'ISOSOMAL UNASE II)	WS08d01x1 NC_CGAP_KId11 Home saplens culva cione IMAGEZASG017 3 similar to contants micra	Williams Welliams DNA phomosecure 4 contin fromment No. 16	AMERICAPISM INITIAL MACE AS HOME SERVICES CONTROLLED TO THE MACE SERVICES OF	0014300UT INIT MOCOULINE SELECTION SERVICE SELECTIONS FOR MACE SELECTION SERVICE SERVICE SELECTION SERVICE SELECTION SERVICE SELECTION SERVICE	0018004084 INIT MACC OF INITIO SEQUENCE CATA CIGIN INTO CATA INTO CATA CATA CATA CATA CATA CATA CATA CA	QVZ-N I UU46-100500-5 10-600 N I OU46 I IUINO Seprenta COLICA	Chlamydophila preumdniae Artse, section et au evolutione generie.	an32c04.y5 Gessler Wilms tumor Homo septens cunA cione invade: 17 003co o	Drosophila melanogaster tyrosine kinase p45 isotorm (fer) mirkny, complete cus	EST364414 MAGE resequences, MAGB Homo septens cDNA	Homo sapiens chromosome 21 segment HS21 C047	601490280F1 NIH_MGC_69 Homo sepiens cDNA clone IMAGE:3892842 6	601070219F1 NIH_MGC_12 Home saplens cDNA clone IMAGE:3456365 5	601070219F1 NIH_MGC_12 Home saplens cDNA clone IMAGE:3456366 6	
Top Hit Detabase Source				EST HOMAN		EST_HUMAN		EST HUMAN		SWISSPROT		EST_HUMAN	EST_HUMAN	_				EST_HUMAN		SWISSPROT		EST DOMAN		EST_HUMAN	EST HOMAN	EST HOMAN		EST HUMAN	<u> </u>	EST_HUMAN	1	EST_HUMAN	EST_HUMAN	EST HUMAN	
	발	į			틱	ES	뉟	ES.	E	<u>8</u>	눌	ES	8	호	-	킥	퇵	<u>ස</u>	Z	જે	<u> </u>	3	Z			띄	Z	Ü	NT	ŭ.	N	ŭ	ŭ	Ü	1
Top Hit Acession No.	TN 8578676	2004404	1.15-01 302410.1	F03285.1	6763231 NT	1.1E-01 BE393186.1	1.1E-01 X62135.1	1.1E-01 R98948.1	Y07695.1	P97384	X52708.1	AW819412.1	AW819412.1	AF157066.1		1.1E-01 S44857.1		AW0265	8923317 NT	1.0E-01 O62855		1.0E-01 AIB85498.1	1.0E-01 AL161504.2	BF033991.1	1.0E-01 BF239818.1	BF365703.1	1.0E-01 AE002285.2	1.0E-01 AI792349.1	1.0E-01 U50450.1	AW952344.1	1.0E-01 AL163247.2	BE881568.1	BE54554.1	BE545554.1	
Most Similar (Top) Hit BLAST E Value	4.4E-04	1	1.16-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01	1.1E-01 Y07	1.1E-01	1.1E-01 X52	1.1E-01 AW	1.1E-01 AW	1.1E-01 AF1		1.15-01				1.0E-01															
Expression	4 24		٦.٥/	. 0.83	1.57	2.75	1.5	1.28	0.8	0.84	4.1	1,01	1.01	86.6		0.96	1.09	98'0	1	3.86					0.87	2.47	1.88	1.85							
ORF SEQ ID NO:	1		12820	12895		13364	13397	49.440			13639		13976				14652							13468	13649	13858	L		14550		L	L	42784	1	
SEQ ID	7027	2	7803	7981	8280	1			8509			1_	1_	L	1		9670	9837					6366	8442	8643	8850	9275	L	1_	L	L	L			
Probe SEQ ID NO:	976	8	2782	2963	3267	3338	3369	2777	3504	3619	3627	3994	3004	4131		4505	4685	4858	5002	1182		1253	1369	3434	3637	3848	4283	4423	4574	47.67	195	5022	3 2	707	2/14

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Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo espiens nerrenth III dinks associated and	O software House of the land o	Control of the Contro	pages calous reduction and granding and general (LDOX) mRNA, LDOX-2 affete, complete cds	Lectus prince in medians beta-tubulin micha, complete cds	Alse arbanescens mRNA for NADD mile	Home contains the ball of the contains the contracts of the contracts of the contains the contai	name septicine includiasi grown ractor receptor 3 (achondroplasia, thanatophoric dwarfiam) (FGFR3) mRNA	CELL SIDEROE ASS ANTICEN PERSONS (SIDEROE)	0247d11 x1 Sparse NHHMD: S1 Line (GLYCOPROTEIN A33)	2247411 of Some Night B. Sa U.	Profess mirabilis fimbial comes and triangles and a constant const	EST378303 MAGE PROGRESS WAS INC.	RCE-BT0254-031000-041-03 BT0254 U	CM2-BN0023-05020-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0	Veneration esculentim polymeter in the sapients along	602150882F1 NIH MCC 81 Home senton CNA ALL 111 CT CALL	M.capricolum DNA for CONTIG MONTA	Homo sapiens BAIT-associated protein 3 (BA14Da) DAIA	Homo saplens nasopharyngael enithellum specific ambits 4 (NIESO4)	602133088F1 NIH MGC 81 Home saniers china lake General Appendix Proposed Et	601286082F1 NIH MGC 44 Homo sapiens CDNA clone IMAGE 3607863 F	801286082F1 NIH_MGC_44 Homo septens cDNA clone IMAGE:3607653 5	AV732224 HTF Homo sapiens cDNA clone HTFAUA08 5'	Mailuscum contagiosum virus subtype 1, complete genome	Molluscum contagiosum virus subtype 1, complete genome	Molluscum contegiosum virus subtype 1. complete genome	yg88f07.r1 Soares trifant brain 1NIB Homo series cDNA close MADE 114649 Et	MAJOR EPIDIDYMIS SPECIFIC PROTEIN F4 (FPIDIDYMAI DECTEIN BE 201	m/9e01.s1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE-226138 a	Mus musculus pre T-cell antigen receptor abha (Picra), mRNA	Human herpesvirus 1 strain KOS-63, latency-associated transcript promoter parties	600944365F1 NIH_MGC_17 Hamo saplens cDNA clone IMAGE:2860176 6
99001 1100	Top Hit Detabase Source								EST HIMAN	SWISSBOT	EST HUMAN	EST HIMAN		EST HIMAN	EST HUMAN	EST HUMAN		EST HUMAN				EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN				Г	SWISSPROT	EST_HUMAN			EST_HUMAN
1	<u>s</u>	Ξ	2	1	15	2	Ę	4503710 NT	:   <u> </u>		I S	I	Ę	T <sub>E</sub>			Ę	183	Ż	4809280 NT	6912525 NT	S	S	舀	<u></u>	Z	N	본	8	S	S	6755216 NT	Σ	<u> </u>
	Top Hit Acession No.	AF099810.1	ıx		ı۹	ाव			E16886	099795	9.6E-02 Al080721.1	A1080721.1	9.6E-02 Z32686.2	9.6E-02 AW986230.1	9.6E-02 BE061729.1	9.5E-02 AW992395.1	9.5E-02 U63374.1	9.4E-02 BF671063.1	233059.1	48097		3F675511.1	9.3E-02 BE391943.1	9.3E-02 BE391943.1	9.3E-02 AV732224.1	J60315.1	J60315.1	J60315.1	354158.1	128631	9.2E-02 AA534354.1	67652	192048.1	3E299722.1
	Most Similar (Top) Hit BLAST E Value	9.9E-02	9.8E-02	9.8E-02	9.8E-02	9.8E-02	9.7E-02	9.7E-02	9.7E-02 B	9.7E-02	9.6E-02	9.6E-02	9.6E-02	9.6E-02	9.6E-02	9.5E-02/	9.5E-02	9.4E-02	9.4E-02	9.3E-02	9.3E-02	9.3E-02 B	9.3E-02	9.3E-02 I	9.3E-02/	9.2E-02 U60315.1	9.2E-02 U60315.1	9.2E-02 U(	9.2E-02 R54158.1	9.2E-02 Q28631	9.2E-02 ∤	9.2E-02	9.2E-02 U92048.1	9.2E-02 BI
	Expression Signal	1.36	1.41	4.03	5.22	5.22	1.38	1.11	4.1	3.83	0.92	0.92	5.54	1.16	76.0	2.27	0.93	2.69	5.14	1.7	6.91	2.33	3.03	3.03	231	7.78	7.76	7.76	3.57	3.85	0.82	1.08	0.98	0.72
	ORF SEQ ID NO:	13229		13101	14081	14082	11379		12300		12054	12055	14191	14800	14926	13969	14902	11877	13796			13222	14012	14013		10287	10298	10289		13140	13269			1
	Exan SEQ ID NO:	8208	5583	8088	9606	9606	6330	6547	7177	8876	6951	6951	9212	9825	8948	8983	8924	6788	8791	7939	7978	8188	9025	8025	3	20	5291	5291	7145	<u>8</u>	8248	8513	9110	9174
	Probe SEQ ID NO:	3192	228	3072	4102	4102	1332	1550	2188	3875	1966	1986	4219	4842	4972	3985	4947	1797	3788	2820	<b>5829</b>	3182	4028	4029	CR C	97	87	877	2168	3106	3233	3505	4118	4181

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Table 4
Single Exon Probes Expressed in HBL100 Cells

Top Hit Descriptor  Counted las Mis-CK gene  O. cunted las Kis-CK gene  FRECEPTOR, ADVL17 (ADVLI-F PECURSOR) (FR-ALPHA) (FOLATE RECEPTOR 1) (FOLATE  RECEPTOR, ADVL17) (ADVLI-F PECURSOR) (FR-ALPHA) (FOLATE RECEPTOR 1) (FOLATE  RECEPTOR, ADVL17) (ADVLI-F PECURSOR) (FR-ALPHA) (FOLATE RECEPTOR 1) (FOLATE  ANTIGEN MOV18) (KB CELLS FBP)  MNS991 bar NOL CGAP_LU24 Home septens a DNA clone INAGE-3178942 3' stimilar to contains Alutery 18 (KB CELLS FBP)  MNS991 bar NOL CGAP_LU24 Home septens a DNA clone INAGE-3178942 3' stimilar to contains Alutery 18 (KB CELLS FBP)  MNS991 bar NOL CGAP_LU24 Home septens a DNA clone INAGE-3178942 3' stimilar to contains Alutery 18 (KB CELLS FBP)  MNS991 bar NOL CGAP_LU24 Home septens a DNA clone INAGE-328936 complete a contains Alutery 18 (KB CELLS FBP)  MNS991 bar NOL CGAP LU24 Home septens a DNA clone INAGE-328936 for complete a contains Alutery 18 (KB CELLS FBP)  MNS991 bar NOL CGAP Home septens a DNA clone INAGE-328936 for contains Alutery 18 (KB CELLS) (KB CELL	mussurus JNN maracing protein-sa (Jip3) mRNA, complete cds Methanobacterium thermoautotrophicum from bases 1176181 to 1189408 (section 101 of 148) of the complete genome
Ggallus Mia-CK gene O. cuniculus k12 karatin gene HUMAN Arabidopsis thaliana DNA chromosome 4, confectors thaliana DNA chromosome 6, confectors thaliana decorption of the chromosome 6, confectors thaliana globulin (Salmin science) Dichyostelium discoldeum spore cost structural confectors thaliana globulin (Salmin science) Dichyostelium discoldeum spore cost structural confectors thaliana globulin (Salmin science) Dichyostelium discoldeum spore a gene confectors thaliana dichyome appiens characters and plasma membrane ce Homo sepiens zinc finger protein 92 (ZFP92), e genes, complete cds; and plasma membrane ce Mare and plasma membrane ce deservations and plasma membrane ce deser	Methanobacterium thermo complete genome
TOP HIT Detabese Source NT	Ż
Acession 0 252 252 422 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	
Most Similar (Top) Hit BLAST E Value 9.2E-02 X98402.1 9.1E-02 X77865.1 9.1E-02 AV477256 9.0E-02 P15328 9.0E-02 AF279132 9.0E-02 AF279132 9.0E-02 AF279132 9.0E-02 BF701583 9.0E-02 BF701583 9.0E-02 BF701583 9.0E-02 BF701583 8.9E-02 BF701583 8.9E-02 BF701583 8.9E-02 BF701583 8.9E-02 AA299128 8.8E-02 C00288 8.7E-02 U82695.2 8.7E-02 U82695.2	8.7E-02 AE
Signal 1.99 1.99 1.99 1.99 1.99 1.99 1.99 1.9	1.08
14472 10063 14472 10063 14274 14158 14166 14167 1467 1	
Exan SEQ ID NO:	9066
Probe SEQ ID NO: 4604 4504 4504 4504 4504 4504 4504 4504	4929

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo sapiens Xq pseudoautoscmal region; segment 2/2	601304016F1 NIH_MGC_21 Hamp sepiens aDNA clane IMAGE:3638643 5	Trichomonas vaginalis beta-tubulin (btub1) gene, complete cds	Dictyostelium discoldeum adenylyl cyclase (acrA) gene, complete cds	Helicobacter pylori 26695 section 130 of 134 of the complete genome	zd44e11.r1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone IMAGE:343532 6*	Cavia porcellus glycoprotein alpha-subunit mRNA, complete cds	Cavia porcellus glycoprotein alpha-subunit mRNA, complete cds	HYPOTHETICAL LIPOPROTEIN MG309 HOMOLOG PRECURSOR	Homo sepiens gene for fukuitin, complete cds	Gallus gallus mRNA for for OBCAM protein gamma tsoform	Canis familiaris glutamate transporter (EAAT4) mRNA, complete cds	Homo sapiens chromosome 21 segment HS210008	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 10	Homo saplens chromosome 21 segment HS21C008	LEUCOCYTE ANTIGEN CD87 PRECURSOR	LEUCOCYTE ANTIGEN CD87 PRECURSOR	LEUCOCYTE ANTIGEN CD97 PRECURSOR	Mus musculus zinc transporter (ZnT-3) gene, complete ads	AU119830 HEMBA1 Hamo saplens aDNA done HEMBA1006744 6	Pseudomonas putida malonate decarboxylase gene cluster (mdcA, mdcB, mdcC, mdcD, mdcE, mdcG,	mdcH, mdcL and mdcM genes), complete cds	Pseudomonas aeruginosa PA01, section 234 of 529 of the complete genome	EST368723 MAGE resequences, MAGC Homo septions curva	Human gene for dihydrolipoamide succinytransferase, complete cos (exon 1-15)	Human gene for dihydrolipoamide succinylfransferase, complete cas (exon 1-10)	PM3-BT0347-170200-001-b08 BT0347 Homo sapiens CUNA	601855548F1 NIH_MGC_67 Home sapiens cDNA clone IMAGE:4076619 6	Thermoplasma acidophilum complete genome; segment 5/5	EST378191 MAGE resequences, MAGI Homo sapiens cultiva	Homo sapiens cAMP responsive element binding protein-like 2 (CREBL2) mrtvA	ti31g02.x1 NCI_CGAP_Gas4 Hamo saplens cDNA cone IMAGE:2132114 3	M.musculus gene for gelatinase B	Molluscum conteglosum virus subype 1, compress gardine
3000	Top Hit Database Source		EST_HUMAN				EST_HUMAN			SWISSPROT							SWISSPROT	SWISSPROT	SWISSPROT		EST_HUMAN				L HOWAN			EST_HUMAN	EST_HUMAN		EST_HUMAN		EST_HUMAN		
{		호	EST	¥	L	IN	EST	N	N	SW	Ł	Ż	IN	Ŋ	Ł	N	NS.	NS.	δĶ	<u>TN</u>	ES	_	뉟	뉠	입	호	뉟	S	ES	Ż	ES	A N	ES	뉟	불
Diffino	Top Hit Acession No.	4,1271736.1	8.6E-02 BE408667.1	8.6E-02 L05468.1	8.6E-02 AF153362.1	8.5E-02 AE000652.1	W69330.1	AF257213.1	AF257213.1	P76334	8.3E-02 AB038490.1	8.2E-02 Y08170.2	8.2E-02 AF167077.2	8.2E-02 AL163208.2	8.2E-02 AL161498.2	8.2E-02 AL163208.2	P48960	P48960	P48960	U76009.1	8.2E-02 AU119830.1		8.1E-02 AB017138.1	AE004873.1	AW954653.1	D26535.1	8.0E-02 D26535.1	8.0E-02 BE067219.1	8.0E-02 BF246744.1	8.0E-02 AL445067.1	8.0E-02 AW966118.1	4503034 NT	A1434202.1	8.0E-02 X72794.1	U60315.1
	Most Similar (Top) Hit BLAST E Value	8.6E-02 AJ	8.6E-02	8.6E-02	8.6E-02	8.5E-02	8:4E-02 W	8.4E-02 AF	8.4E-02 AF	8.3E-02	8.3E-02	8.2E-02	8.2E-02	8.2E-02	8.2E-02	8.2E-02	8.2E-02	8.2E-02 P4	8.2E-02 P48960	8.2E-02	8.2E-02					8.0E-02									
	Expression Signed	822	1.3	3.35	3.97	1.85	3.69	0.99	66'0	5.85	0.83	6.17	2.51	2.44	1.37	1.16	5.75	5.76	6.75	3.34	1.29		1.08	1.05	4.12	9.29	9.29		2.75		0.73				
	ORF SEQ ID NO:	11276	12281	13148		12428	12674	14202	14203	13532	14532		11521			13889	14139						11520	14999	10072	11728		L	L	12869			14807		10971
	Egan SEQ ID NO:	L	7161	L	8288	7307	7778	9222	9222	8520	9547	6357					L	L	L	L	L	L	6461	10031	7711	7752		6851		L		L			5839
	Probe SEQ ID NO:	1233	2182	3114	3559	2333	2596	4228	4228	3512	4559	1360	1465	3000	3713	3901	4161	4161	4161	4909	5027		1464	5062		1659	1659	1862	2400	2830	3726	3958	4631	4669	5060

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Table 4
Single Exon Probes Expressed in HBL100 Cells

		<b>,</b>	_	_	_			_		_	·	T -	_		_		P	T	4		5		L	Ч			74	7	إذ	<b>5</b> ,
Top Hit Descriptor	600943191F1 NIH_MGC_15 Hamp saplens cDNA done IMAGE:2959510 5	ar88c08.x1 Berstead colon HPLRB7 Home septens cDNA clone IMAGE:2173846 3' straiter to gb:228878 80S RIBOSOMAL PROTEIN L38 (HUMAN);	Mus musculus actory etimulating factor 1 receptor (Ceffr), mRNA	Mus musculus colony stimulating factor 1 receptor (Csf1r), mRNA	Arabidopsis thaliana RXW241, mRNA, partial cds	0059402.y5 NCI_CGAP_Lu5 Homo sepiens cDNA done IIAAGE:1570467 6' similar to contains L1.t3 L1 repetitive element;	0059402.y5 NCI_CGAP_Lu5 Homo septens cDNA done IMAGE:1570467 5' similar to contains L1.33 L1 receitline demant:	Sus scrofa telomerasa RNA pseudocene	Sus scrofa telomerase RNA pseudopane	600943055F1 NIH MGC 15 Home septems cDNA clone IMAGE 2959603 for	800943055F1 NIH MGC 15 Hamo sapiens cDNA clane IMAGE:2959683 5	tg48g12.x1 Soeres_NFL_T_GBC_S1 Homo septens cDNA clone IMAGE:2112070 3' similar to contains	WENTUCK MENTO REPORTED	Homo sapiens partial AF-4 gene, excns 2 to 7 and Alu repeat elements	601316428F1 NIH_MGC_8 Homo septens cDNA clone IMAGE:3834903 6	EST112214 Cerebellum II Homo sapions cDNA 5' end similar to similar to protocardherin 43	Homo septens solute center family 6 (neurotransmitter transporter, glycine), member 9 (SLC6A9), mRNA	Homo sapiens solute carrier family 6 (neurotransmitter transporter, glycine), member 9 (SLC6A9), mRNA	Homo saplens chromosome 21 segment HS21C078	Homo sapiens IL-18 gene for interleukin-18, intron 1 and exon 2	RC5-LT0054-260100-011-H09 LT0054 Homo sapiens cDNA	wf43h01.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA cione IMAGE:23583853'	Homo saplens ADP/ATP carrier protein (ANT-2) gene, complete cds	Rattus norvegicus Activin receptor like kinase 1 (Acvrl1), mRNA	Mus musculus ubiquintin o-terminal hydrolase related polypeptide (Uchrp), mRNA	601658738R1 NIH_MGC_69 Homo sapiens cDNA clone IMAGE:3886209 3'	601658738R1 NIH_MGC_69 Homo sapiens cDNA clone IMAGE:3886209 3'	Thermotoga maritima section 101 of 138 of the complete genome	CMO-NN1004-130300-284-g08 NN1004 Homo sepiens cDNA	
Top Hit Database Source	EST_HUMAN	EST HUMAN				EST HUMAN	EST HIMAN			EST HIMAN	EST HUMAN		ESI JONAN		EST_HUMAN	EST_HUMAN					EST_HUMAN	EST_HUMAN				EST_HUMAN	HUMAN		EST_HUMAN	
	ESI	EST	뉟	뉟	눌	EST	I SH	Ę	Ę	EST	EST		3	뒬	EST	EST	支	Ę	Ę	Ł	EST	EST	Ę	N	Ę	EST	_	Ž	ESI	
Top Hit Acession No.	7.9E-02 BE250008.1	7.9E-02 Al582029.1	6881044 NT	6881044 NT	7.9E-02 AB008019.1	7.8E-02 AI793276.1	7 8F-02 A1783278 1	7.8E-02 AF221942.1	7.8E-02 AF221942.1	7 8E-02 BE250048 1	7.8E-02 BE250048.1	7	AI418520.1	7.7E-02 AJ238093.1	7.6E-02 BE514432.1	7.6E-02 AA296447.1	5802083 NT	5902083 NT	6327	7.5E-02 AB015961.1	AW838547.1	7.4E-02 AI807885.1	L78810.1	6978442 NT	6678492 NT	7.3E-02 BE964961.2	7.3E-02 BE964961.2	7.3E-02 AE001789.1	AW900281.1	
Most Similar (Top) Hit BLAST E Velue	7.9E-02	7.9E-02	7.9E-02	7.9E-02	7.9E-02	7.8E-02	7.85-02	7.8E-02	7.8E-02	7.85-02	7.8E-02	100	1.8E-02 AI41	7.7E-02	7.6E-02	7.6E-02	7.5E-02	7.5E-02	7.5E-02 AL1	7.5E-02	7.4E-02 AW	7.4E-02	7.4E-02	7.4E-02	7.4E-02	7.3E-02	7.3E-02	7.3E-02	7.3E-02 AW	
Eppression	2.64	11.99	5.05	5.05	1.4	1.59	60	-	-	1.25	3.15		7.02	2.61	2.57	0.82	1.86	1.86	1.15	0.92	1.1	0.77	1.18	2.97	1.75	1.23	1.23	3.86	2.33	
ORF SEQ ID NO:	12203	12948	13763	13764		11228		12426	12427				12002			13354	10826	10827		14348	10523	13633	14541	14628	14752	10514		10709	11508	
SEQ ID NO:	7089	7929	<u>.</u>		9651	6192	6192		L	L	L	1				8334	6798	5798		8368	5510	8522	9554		89/6	5503	6503	10/9	7748	
Probe SEQ ID NO:	2109	2910	3760	3760	4666	1191	1191	2332	2332	3663	4914	900	8	3506	3303	3324	776	776	1877	4377	474	3514	4566	4650	4784	466	466	876	1450	

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo sapiens chromosome 21 segment HS21C102	Mus musculus transcription factor USF2 (USF2) gene, exons 8-10 and complete cds	Methanobacterium thermoautotrophicum from bases 1029155 to 1039934 (section 88 of 148) of the complete	Methodochected in the mountainshiring from bases 1029455 to 1039924 (section 88 of 148) of the complete	perome	Homo sapiens chromosome 21 segment HS21C101	Homo saplens chromosome 21 segment HS21C101	Human immunodeficiency virus type 1 isolate 28 reverse transcriptase (pci) gene, internal fragment, pertial ods	UI-H-BW0-41-4-05-0-UI.s1 NCI_CGAP_Sub6 Homo suplens aDNA alone IMAGE:2732049 3'	602077767F1 NIH_MGC_62 Homo septens cDNA clone IMAGE:4251950 6	Human Immunodeficiency virus type 1 (D9) proviral structural capaid profein (gag) gene, partial eds	Pseudomonas aeruginosa PA01, section 451 of 529 of the complete genome	601872281F1 NIH_MGC_63 Hamo saplens cDNA clane IMAGE:4092981 51	ha10b06.71 NIH_MGC_7 Homo sepiens cDNA clone IMAGE:2823921 5' similar to gb:X62851_me1 PEPTIDYL-PROLYL CIS-TRANS ISOMERASE A (HUMAN); gb:X52803 Mouse mRNA for cyclophilin	(MOUSE);	COLLAGEN ALPHA 1(XVI) CHAIN PRECURSOR	Martiellia Mout-1 gene	266f04,s1 Strategene colon (#837204) Homo sepiens cDNA clone IMAGE:608599 3'	ULH-BIT-ecy-c-07-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2716020 3:	ai65a12.s1 Soares_testis_NHT Homo sapiens cDNA clone 1376678 3' similar to gb:K03002 60S RIBOSOMAL PROTEIN L32 (HUMAN);	QV4-BT0407-280100-090-e10 BT0407 Homo saplens cDNA	CM0-UM0001-060300-270-e12 UM0001 Homo sapiens cDNA	Canis familiaris inducible nitric code synthase mRNA, complete cds	601816291F1 NIH_MGC_56 Homo saplens cDNA clone IMAGE:4050071 5	Homo sapiens chromosome 21 segment HS21C010	Homo sapiens chromosome 21 segment HS21C010	Homo septens regulator of Gz-selective protein signaling (ZGAP1) mRNA, and translated products	26S PROTEASOME REGULATORY SUBUNIT 63 (NUCLEAR ANTIGEN 2107)	
	Top Hit Detabese Source									EST_HUMAN	EST_HUMAN			EST_HUMAN		EST_HUMAN	SWISSPROT		EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN		T_HUMAN				SWISSPROT	- :
		Ę	Ę	Ė		노	E	호	뒫	EST	EST	NT	뉟	ESI		ES	MS	Ľ	ESI	ES	ES.	ES	ES.	ĮN.	EST	IN	Z	3 NT	NS	
,	Top Hit Acession No.	7.3E-02 AL163302.2	7.3E-02 U12283.1	A E. COOD 60 4	7.4E-V2 AEWUDD2.1	AE000882.1	7.2E-02 AL183301.2	7.2E-02 AL163301.2	U14784.1	AW298322.1	7.2E-02 BF572307.1	7.1E-02 L02290.1	AE004890.1	7.1E-02 BF208802.1		BE208576.1	Q07092	7.0E-02 X96677.1	7.0E-02 AA056343.1	AW138152.1	7.0E-02 AA815438.1	7.0E-02 BE070264.1	7.0E-02 AW792962.1	AF077821.1	7.0E-02 BF381987.1	AL163210.2	6.9E-02 AL163210.2	4507968 NT	6.9E-02 Q06364	
	Most Similar (Top) Hit BLAST E Value	7.3E-02	7.3E-02	7 20 00	1.4E-02	7.2E-02	7.2E-02	7.2E-02	7.2E-02 U14	7.2E-02	7.2E-02	7.1E-02	7.1E-02	7.1E-02		7.1E-02 BE:				7.0E-02 AW					L					
	Expression Signal	11.5	1.14		7.	1.2	1.67	1.67	1.86	1.62	6.17	1.58	1.06	5.78	16.	1.09	1.16	0.92	1.17		0.82		96'0	1.2		17.58				
	ORF SEQ ID NO:				10202	10203		11502		13799	14192	11940		12322	<u> </u>	14946	10562	l	11797	12993	13810		L	14076		10548		<u> </u>	13709	
İ	SEQ ID NO:	7756	}_		300	6190	L				L		1_		<u> </u>	8968				L	8805		L					1	L	1
	Profe SEQ ID NO:	1808	4838		120	120	1448	1446	2478	3791	4220	1863	2225	2230		4897	524	1470	1724	2960	3802	3931	4017	4093	4773	510	510	1313	3703	

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Acession Top Hit Database Top Hit Descriptor	28S PROTEASOME REGULATORY SHIPLINT SYAM IS: EAD ANTIGEN AGE	ae30f02.rt Gessier Wilms turner Homo septems cDNA clane IMAGE:897339 6' similar to gb:M22382	4830f02.rf Gessler Wilms turner Home septens CDNA clone IMAGE:897339 6' similar to gb:M22382	HOMO SENIORS CHICALE HOLD THE PROJECT CHICAGON,	ai75an6 st Shares tootic MILT U.	AITEM STATES TO States the MILTITUTE TO SECURE ALIVE AND STATES STATES AND ST	BITSB08 st Scarce testis NIT I name septens clone 1376626 3'	Oncombrothis myles TAB4	19979604.x1 Source NF T CRC St Home College Co	HOMEOBOX PROTEIN HOX DA (CHOX A)	att 2609.xt Barstsad acrta HPLRB Hono saplens cDNA clone IMAGE:2354920 3' similar to SW-1 NA NYCCO Description	Discaphile majorate and pure - Diff	Mus misculine Canal 2	VIRBAIO 81 Source placests Nicolate 11.	Homo septens maschiella (A.K.) A.V.	Homo seriens mescribain (MSLN), denscribain (M	Homo sablens TESTIN 2 and TESTIN 3 and 1 a	INTER-ALPHA-TRYPSIN INHIBITOR HEAVY CHAIN IS DEEX IESSO, ITT. 1-71.00	INTER-ALPHA-TRYPSIN INHIBITOR HEAVY CHAIN H2 PRECIDENCE (ITI HEAVY CHAIN HZ)	601671046F1 NIH_MGC_20 Homo sepiens cDNA clone IMAGE-3054178 F	Homo saplens E2F-like protein (LOC51270), mRNA	Xenopus laevis alpha(E)-catenin mRNA, complete cds	Aquifex sections section 98 of 109 of the complete genome.	A.carterae precursor of peridinin-chlorophyla-grotein (PCP) gene	Themotoga martima section 89 of 136 of the complete	Thermotoga martima section 89 of 136 of the complete general	Mus musculus histore descemase 5 (Hder5) mRNA	Homo sapiens chromosome 21 segment HS21C047	Mus musculus major histocompatibility locus class III regions Heo70t gene, partial ods; smRNP, G7A, NG23, MMS homolog, CLCP, NG24, NG28, and NG28 never consider of the consideration of the considera	SPUBB WILLIAMS AND THE STATE A
Top Hit Database Source	SWISSPROT	EST HUMAN	- FRE	L	EST HUMAN	EST HUMAN	EST HUMAN	L	EST HUMAN	SWISSPROT	- FR	IN IN	L	EST HIMAN	-	Ę	N	SWISSPROT	SWISSPROT	EST_HUMAN	<u> </u>	NT	N	Į,	본	N <sub>T</sub>	H	NT	Ŋ	
Top Hit Acession No.	206364	AA496759.1	AA496759.1		Γ	Ī	Ī	Ī		P17278	Al735509.1			T	7108357	7108357 NT	4F260225.1			3F027639.1	7706068 NT		4.1		NE001777.1	VE001777.1	TN 6299692	L163247.2 N	F109905.1	1
Most Similar (Top) Hit BLAST E Value	6.9E-02	6.8E-02	6.8E-02	6.8E-02	8.8E-02	8.8E-02	6.8E-02	8.7E-02	6.7E-02	6.7E-02	6.6E-02	6.6E-02	6.6E-02	6.6E-02	6.6E-02	6.6E-02	6.6E-02			6.5E-02	6.5E-02	8.5E-02	6.5E-02 /	6.4E-02)	6.4E-02	6.4E-02	6.4E-02	6.4E-02	6.3E-02 A	
Expression Signal	1.03	1.06	1.06	3.07	1.01	1.01	1.01	1.66	1.32	4.58	1.14	96.0	1.74	9.7	2.63	2.63	1.53	9.95	9.95	7	3.15	2.48	1.65	1.62	0.96	96.0	1.88	1.21	2.39	
ORF SEQ ID NO:	13710	11820	11921				13055			13646	11376	11399	12210	13413			13953	14783	14784	10608	11022	11414	11768	10801	11765	11766	12984	14883	11788	
Exan SEQ ID NO:	8707	6833			8044	8044	8044	6495	8843	8641	6328	6349	7097	8390	8406	8406	888	9803	883	800	808		2600	2603	9690	0698 8	7965	9905	6711	
Probe SEQ ID NO:	3703	1843	1843	1865	3027	3027	3027	1497	1854	200	1330	1352	2117	3382	3398	3398	3965	4819	4819		7/8	200		2	<u>288</u>	1895	2946	\$2g	1716	

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Single Exon Probes Expressed in HBL100 Cells

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	Ten Life Decombation	milipoper w. da.	LEAT SUCK BEATTING	Application and a supplied of the supplied of	Rattus novvegicus differentation-associated Na-densentant incommis-	complete eds	52 KD RO PROTEIN (SJOGREN SYNDROME TYPE A ANTIGEN (SS. ANVBOGE ANVERSE)	1997a12.s1 Source Infant brain 1NIIB Homo sapiens cDNA clone IMAGE:41477 3' similar to ab:X67198 odar	INVANCALITION ELONGATION FACTOR S-II (HUMAN);	numen mrNvA, Ad terminal portion	Arabidopsis thaliana K+ Inward rectifying channel protein (Aff(C1) gene, complete cds	S. Schola m.R.N.A. for Man9-mannosidase	4gaugeus XI Sogres_NFL_T_GBC_S1 Hamo septens cDNA clane IMAGE:1842470 3'	I hermotoga maritima section 89 of 136 of the complete genome Mesocestoldes corti mitochondrial DNA, NADH dehydrogenese subunit 4. fRNA_cin. fRNA_bese subunit 4. fRNA_cin.	A I Pase subunit 6, and NADH dehydrogenase subunit 2	20/8c04.r1 Strategene Hel a cell s3 937216 Homo sapiens cDNA clone IMAGE:628310 6	ZP/0004.11 Strategene HeLe cell s3 837216 Homo sapiens cDNA clone IMAGE:628310 6	EST 84205 Colon adenocarcinoma IV Homo sapiens cDNA 5' end similar to tissue-specific protein	ES 184266 Colon adenocarcinoma IV Homo saplens cDNA 5' end similar to tissue-specific profein	601658150R1 NIH_MGC_68 Homo septens cDNA clone IMAGE:3876060 3'	rc-1-D10001-290100-012-e10 D10001 Home sepiens cDNA	Mus musculus pos tumor suppressor gene, exon 10 and 11, partial cds; alternatively spliced	KINEON DEAVY STAND AND SERVED BY THE SERVED OF THE SERVED	Populus Hoberton (CAACHY)	Nemododa marilina section 97 of 43e of th	W24c02 X NCI COAD KIALA LA COMPANIA GENOME	w24c02x1 NCI CGAP Kid11 Home series cDNA close IMAGE:2544578 3'	qh8601.x1 Soares_feta_liver_spleen_INFLS_S1 Home saplens CDNA clone IMAGE:1848697 9' similar to	4h56f01xf Soeres fetal Iver spleen 1NFLS_S1 Home saplens cDNA clone IMAGE:1848697 3' similar to			riomo sapiens qual adaptor of phosphotyrosine and 3-phosphoinositides (DAPP1), mRNA	
	Top Hit Database	Source	TOGGSSIWS			NT.	SWISSPROT	EST HIMAN	TOWOL TO	E L	111	ECT CIPANI	NAMOU TO I	_	-	EST HUMAN	EST HOMAN	EST HUMAIN	EST HUMAN	HUMAN	THE PERSON AND THE PE		TOGGSON			EST HUMAN	Т			יישומויי			
	Top Hit Acession			22		6.2E-02 AF271235.1			Ī	Ī	2503.4	Τ	8 OF-02 AF004 777 4	Τ	T	ŀ	Т	1	1	5.9E-02 AW934719 4	T	T	T	Ī	Γ	Γ	5.8E-02 AW051827.1 ES			T	100g	NIGOTO	
Most Cimilar	(Top) Hit BLAST E	Value	6.3E-02 P37092	6.2E-02		6.25-02/	6.2E-02	6.2E-02 F	6.1E-02 D18471 4	6.1E-02 U73325.4	8.1E-02 Y1	6.1E-02 A	8 OF 02 A	A OF O2 A	8 OF 02 A	6.0E-02	6 OF 02 A	6.0E-02	8 OF 02 B	5.9E-02	5.9E-02 A	5.8E-02 D90110.1	6.8E-02 Q61768	6.8E-02 A	5.8E-02 AI	5.8E-02 A	5.8E-02 A	5.8E-02 Al247505.1	5 8E-02 AI247KN5 1	5.8E-02 A	5.8E-02 78		
	Expression Signal		2.65	3.64		- 3	02.0	.68	3.36	217	0.92	1.37	0.88	2 18	0.0	0.91	1.82	1.62	2.87	6.97	2.79	3.97	2.52	0.94	1.8	5.55	5.55	4.67	4.67	2.28	4.11		•
	ORF SEQ ID NO:			14108				14963	10324		14947		11282		10183	10184	13198	13197		10300	12952		11683		13591	14204	14205	14392	14393	-	14942		
ı	SEQ ID		8530	9124	9205	PAA		9987	5314	8886	9971	6966	6241	7857	6173	6173	8174	8174	8556	5292	7933	5936	6617	7810	8288	8223	8233	9408	9408	9431	8864		
	SEO ID		3523	<del>2</del>	4212	4460		5016	254	3885	5000	5018	1243	2700	2866	2866	3158	3158	3549	229	2914	920	1620	2789	3579	4228	4229	4416	4416	4441	4992		

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	ou83b05.s1 NGI_CGAP_Br2 Homo septens cDNA clone IMAGE:1632465 3' similar to WP:C37A2.2 CE08611;	Hamo sapiens depermine transporter (SLC6A3) gene, complete cds	Chironomus thummi thummi globin VIIA.1 (ctt-7A.1), globin 8.1 (ctt-8.1), globin II-bela (ctt-2bela), non- functional globin XIII (ctt-13RT), globin XII (ctt-12) and globin XI (ctt-11) genes, complete ods	EST378865 MAGE resequences, MAGI Homo sapiens cDNA	Bos taurus lysozyme gene (cow 3), complete cds	Hydrocotyle rotundifolia ribosomal protein L16 (rpi16) gene, Infranc chloroplast gene for chloroplast product	801494578F2 NIH_MGC_70 Home sapiens cDNA clone IMAGE:3896810 5	Lycopersicon esculantum LE-ACS8 mRNA for 1-aminocyclopropane-1-carboxylate synthase, complete cds	zs45c01.s1 NCI_CGAP_GCB1 Hamo sapiens cDNA clone IMAGE:700416 3'	H.saplens gene encoding La autoanigen	Mus musculus SH3 domain protein 1B (Sh3d1B), mRNA	Gallid herpesvirus mRNA fragment	Homo saplens HTRA serine protease (PRSS11) gene, complete cds	Oryza sativa rbbi3-1 gene for putative Bowmen Birk trypsin inhibitor	RC5-BT0559-140200-012-C03 BT0559 Hamo sapiens cDNA	QV0-ST0213-021299-082-e09 ST0213 Homo sepiens cDNA	QV0-ST0213-021289-062-e09 ST0213 Homo saplens cDNA	ye37f12.r1 Stratagene lung (#937210) Homo sepiens cDNA clone IMAGE:119951 5' similar to gb:K01506 HLA CLASS II HISTOCOMPATIBILITY ANTIGEN, DP(1) ALPHA CHAIN (HUMAN);	Pseudomonas putida ttgS gene	Drosophila melanogaster laminin B2 gene, complete cds	Drosophila melanogaster laminin B2 gene, complete cds	Pseudomonas putida ttgS gene	Mus musculus caudal type homeobox-1 (Cdx-1) gene, complete ods	Homo saplens meprin A, alpha (PABA peptide hydrolase) (MEP1A) mRNA	Homo saplens partial LMO1 gene for LIM domain only 1 protein, exon 1	Homo saplens partial LMO1 gene for LIM domain only 1 protein, exon 1	Human steroid hormone receptor Ner-I mRNA, complete cds	Zp647D073 6'	Homo sapiens PBII gene for salivary proline-rich protein P-B, complete cds	
Top Hit Database Source	HUMAN			EST_HUMAN			EST_HUMAN		EST HUMAN						HUMAN	<b>EST_HUMAN</b>	EST_HUMAN.	EST_HUMAN										HOMAN		
	EST	Ę	둗	EST	Ę	Ľ	EST	Ę	EST	Ę	툳	뉟	눈	Ę	EST	EST	EST	EST	ΙN	LΝ	M	ΙN	Ż	토	Ę	NT	Σ	EST	国	
Top Hit Acession No.	6.7E-02 Al081644.1	4F119117.1	AF001292.1	5.7E-02 AW966791.1	5.7E-02 M95099.1	AF094455.1	3E904308.1	5.6E-02 AB013100.1	5.6E-02 AA290599.1	K97869.1	6755501 NT	41561.1	4F157623.1	5.4E-02 AJ277468.1	73468.1	391248.1	391248.1	194759.1	AJ276408.1	5.3E-02 M58417.1	6.3E-02 M58417.1	5.3E-02 AJ276408.1	M80463.1	5031908	5.2E-02 AJ277681.1	5.2E-02 AJ277661.1	5.2E-02 U07132.1	5.1E-02 AL134071.1	AB031740.1	
Most Similar (Top) Hit BLAST E Value	6.7E-02	5.7E-02 AF1	6.7E-02	5.7E-02	5.7E-02	6.6E-02	5.6E-02 BE9	5.6E-02	5.6E-02	6.5E-02 X97	5.6E-02	5.5E-02 L41	5.4E-02 AF1	5.4E-02	5.4E-02	5.3E-02 AW	5.3E-02	5.3E-02 T94	5.3E-02 AJ	5.3E-02	6.3E-02	5.3E-02	5.3E-02	5.2E-02	5.2E-02	5.2E-02	5.2E-02	5.1E-02	5.1E-02	
Expression Signal	1.08	1.11	1.02	2.21	1.05	1.7	1.01	1.56	0.84	4.77	3.85	42.	0.75	0.93	6.65	1.48	1.48	7.24	1.9	0.82	0.82	4.71	9.18	66.99	2.21	2.21	3.13	96.0	0.91	 
ORF SEQ ID NO:	13013	13029		13718		11548		14474	14530	12660	13181	14077				11075	11078	11531	12518	12913	12914	13109	14868		13065	13068	14133			
Exan SEQ ID NO:	8001	8016	9632	8716	9535		7203	9498	9545	7546	8159	6806	6268	7972	10058		6046	6473	2395		1691	8095	9894		8057	2508	9151	7278	9827	
Probe SEQ ID NO:	2983	2998	3625	3712	.4546	1495	2226	4506	4557	2583	3143	4095	1270	2953	3337	1036	1036	1476	2424	2872	2872	3079	4916	2221	3040	3040	4156	2303	4845	

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Table 4

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Single Exon Probes Expressed in HBL100 Cells	Top Hit Descriptor	Mus musculus fatty acid emide hydrolass gene, excn 10	Bacillus subtilis complete genome (section 1 of 21): from 1 to 213080	SALIVARY ACIDIC PROLINE-RICH PHOSPHOPROTEIN 1/2 PRECURSOR (PRP-1/PRP-3) (PRP-2/PRP-4) (PIF-FIPIF-S) (PRP-2/PRP-4) (PIF-FIPIF-S) (PROTEIN A/PROTEIN C) [CONTAINS: PEPTIDE P-C]	Oryctolegus cuniculus UDP-glucuronosyltransferase (UGT2B13) mRNA, complete cds	Mus musculus Uno-51 like kinase 2 (C. elegans) (Ulk2), mRNA	Haemophilus influenzae Rd section 97 of 163 of the complete genome	Aniherasa pernyi period clock protein homolog mRNA, complete cds	Chicken 28-kDa vitamin D-dependent calcium-binding protein (CaBP-28) mRNA, complete ods	Homo sapiens ABCA1 (ABCA1) gene, complete cds	Homo sapiana ABCA1 (ABCA1) gene, complete cds	Zea mays phytoene synthase (Y1) gene, complete cds	ATROPHIN-1 (DENTATORUBRAL-PALLIDOLUYSIAN ATROPHY PROTEIN)	zq48a12.s1 Stratagene hNT neuron (#637233) Homo sapiens CDNA clone IMAGE:632928 S' similar to	contains Alu repetitive element;contains element MSR1 repetitive element;	zi78a03.s1 Soares_testis_NHT Homo sapiens cONA clone IMAGE:728428 3'	zt78a03.51 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:728428 3	xg56g10.x1 NCI_CGAP_Ut4 Home sepiens cDNA clone IMAGE:2632386 3	xg56g10.x1 NCI_CGAP_Ut4 Homo sapiens cDNA cione IMAGE:2632386 3	Human mRNA, Xq teminal portion	Human mRNA, Xq terminal portion	Arabidopsis thaliana AP2 domain containing protein RAP2.7 mRNA, partial cds	zo49b02.s1 Soares_senescent_fibroblasts_NbHSF Homo saplens cDNA clone IMAGE:325611 3' similer to gb:M30838 LUPUS KU AUTOANTIGEN PROTEIN P86 (HUMAN);	Tetrahymena rostrata histone H3II and histone H4II Intergenic DNA	PM0-HT0339-251199-003-g05 HT0339 Homo saplens cDNA	Escherichia coli K-12 MG1655 section 335 of 400 of the complete genome	am50d02.s1 Johnston frontal cortex Homo sepiens cDNA clone IMAGE:1538979.3' similar to TR:P80533	P90533 LIMA ;contains element LTR1 repetitive element ;	AV727059 HTC Homo sapiens cDNA clone HTCBW C01 6	/xn24f03.x1 NC_CGAP_Kid11 Homo sepiens cDNA clone IMAGE:2894863 S' semilet to SW:GKTHUMAAN  Q12849 G-RICH SEQUENCE FACTOR-1 ;	PMO-HT0339-251199-003-005 HT0339 Homo sapiens cDNA	PN/0-HT0339-251199-003-005 HT0339 Homo sapiens cDNA	
n Probes E	Top Hit Database Source			SWISSPROT									SWISSPROT		EST HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN				EST HUMAN		EST HUMAN	1		EST HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	
Š.		호	호	8	눋	TN 0	Z	Ż	Ę	눌	Ł	Ż	NS	-	S	ES	ES	ES	ន្ន	Ż	Ä	Ξ		E	<u> </u>	E	┞	ŭ	E	<u>ü</u>	1	įω	4
Single	Top Hit Acesslon No.	5.0E-02 AF098004.1	289104.1	P02810	U72742.1	7305810 NT	5.0E-02 U32782.1	5.0E-02 U12769.2	4.9E-02 M14230.1	4.9E-02 AF275948.1	4.9E-02 AF275948.1	U32636.1	P54258		4.9E-02 AA188940.1	4.9E-02 AA400914.1	4.9E-02 AA400914.1	4.9E-02 AW167821.1	4.9E-02 AW167821.1	4.8E-02 D16471.1	4.8E-02 D16471.1	AF003100.1	4 RE-02 W61983 1	4.8E-02 X17144.1	BE153583.1			A1014255.1	AV727059.1	AW296028 4	BE183583 1	4.0E-02 DE 163583 1	10-100001
	Most Similar (Top) Hit BLAST E Value	6.0E-02	5.0E-02	5.0E-02 P02810	5.0E-02 U	5.0E-02	5.0E-02	5.0E-02	4.9E-02	4.9E-02	4.9E-02	4.9E-02	4.9E-02		4.9E-02	4.9E-02	4.9E-02	4.9E-02	4.9E-02			1	4.8F.02					4.6E-02	4.6E-02				
	Expression Signal	1.87	12.28	2.86	1.3	1.24	1.04	7.06	30.11	2.47	2.47	0.89	1.52		2.64	0.71	0.71	2.02	2.02	1.13	2.65	8.84	1 27					0.71					
	ORF SEQ ID NO:	10528		12029	11021			13615		10431	10432	12838				13530	13531	14660	14661	10390	10390	10531	13300						11390		1	10330	1
	SEQ ID NO:	5516	6186	08830	5987	8276	8523	8607	6285	5417	5417	7823	8232		8498	8519	8519	7196	9677	5381	5381	L		١				6271	1		1		908/
	Probe SEQ ID NO:	479	1186	194	2748	3263	3515	3600	23	368	368	2803	3217		3480	3511	3511	4692	4692	328	88	485	3.56	8438	268	2	3	1273	4342		8147	2736	3416

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Mus musculus nucleoler RNA helicase IVGu (do.21) gene, complete ode	RETINOIC ACID RECEPTOR BETA (RAR-BETA)	Marburg virus strain M/S. Afficationarinessoligi is/ alocalini virus garo, canipas cas	Marburg virus strain M.S. Africa Johannesourg 1970 of Crown virus strain M.S. Africa John S. Office John S. Africa John S. Afr	HEPATOCYTE NUCLEAR FACTOR 3-BETA (HNF-3B)	Xylella fastidiosa, section 110 of 229 of the Complete genome	Homo sapiens chromosome 21 segment HSZTCU/o	601652154F1 NIFL MGC 82 Floring saparts cults while invocations of	HYPOTHETICAL PROTEIN (ORF 2284)	QV2-PT0012-010300-070-g02 PT0012 Framo equipme s CONP.	Myxococcus xantrus settreturecrine ruisse Friio (pwilo) gene, waipieus cus	Homo sapiens S164 gene, partial cds, PS1 and nyponiencal protein gaines, curippete cus, and climing and control an	per dar cus.	Homo expens of or gans, parties cus, right and remaining the remaining control of gans, parties of s	Months exertilis muceln heary chain FM3A (FM3A) mRNA, complete cds	United analysis chromosome 21 segment HS21C010	FIGURE Separation of the company of	Home septembly promy contents and MAZNAZO00020 5	AUTOS227 NTORMO Homo ganiens cDNA clone NT2RM2000020 5'	ACTESSED NICE COAD BIT Homo seniens CONA clone IMAGE:2545584 3' similar to TR:Q63291 Q63291	WX34901X11NC_CONINTERMEDIATION CONTRIBUTED TO THE PROPERTY CONTRIBUTION OF THE PROPERTY CONTRIBUTION	Thermoplasma acidophilum complete genome, segment 452	TRANSFORMING PROTEIN WAT	80201/105F1 NCL COAP DILECT HALL SAPARE CONTRACT TO THE COMPLETE CONTRACT TO THE	Chiamydra muridai uni, secural co el co el companyone murida de companyone co	CV1-INNUUTZ-18U4UV-194-100 INNUUTZ-18U4UV-194-100 INNUUTZ-18U4UTZ-18U4UTZ-18U4UTZ-18U4UV-18U4UTZ-18U4UTZ-18U4UTZ-18U4UTZ-18U4UTZ-18U4UTZ-18U4UTZ-18U4UTZ-18U4UTZ	L. monocytogenes type 3 per use top yours convious convio	WD98h01.XI NCI COART F128 Italia estrata Color Control Color	Tromo sapirals till to the August of the Supy Homo sapiens con MAGE:3084134 S	OF-TH-BWI HEIMPING CO. TO CO.	TAS ANTICENCOME.	
Top Hit Database Source		SWISSPROT	NT	본	SWISSPROT	NT	N	EST_HUMAN	SWISSPROT	EST HUMAN	NT		Z	5	į.	Ž!	Z	NT.	ESI HUMAN	EST HOMAN	EST_HUMAN	LN	SWISSPROT	EST HUMAN	LN.	EST HUMAN	Į,	EST HUMAN	NT	EST HUMAN	SWISSPROI	
Top Hit Acession No.	4.6E-02 AF220365.1 NT			П									4.4E-02 AF109907.1		T			1	4U123327.1	AU123327.1	AW003645.1	AL445068.1	P23091								P41047	AJ403386.1
Most Similar (Top) Hit BLAST E Value	4.8E-02	4.5E-02	4.5E-02	4.5E-02 /	4.5E-02 P32182	4.5E-02	4.5E-02	4.4E-02	4.4E-02	4.4E-02	4.4E-02		4.4E-02		4.4E-02	4.3E-02				4.2E-02	4.2E-02	4.2E-02	4.2E-02			4.1E-02	4.1E-02					3.9E-02
Expression Signal	98.0	1.67	. 0.81	0.81	4.15	2.17	3.82	3.85	3.33	1.11	1.81		0.99		0.99	6.82	8.44	1.23	1.73	1.9	1.49	2.39	1.61	3.38	1.08		0.67		3.01	3.41		1.85
ORF SEQ ID NO:		10499	11236	11237	11847	12145			<u> </u>	12510	13568		14461				13372		10866		10044		13592					11660	13207		11371	Ш
SEQ ID	000	5481	6200	6200							L	1_	9483			5794	8354	8582	5833	5 5874	5007	$\perp$			Ŀ	L			L			Ш
Probe SEQ ID NO:	4004	44	1189	1199	1768	2051	3640	219	2039	2419	3555		4493		4483	772	3345	3576	812	855	900	168	3580	4600	2605	4347	4064	1603	3170	1103	1326	1921

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Hamo sapiens succinate dehydrogenase complex, subunit C, integral membrane protein, 15tO (SDHC) mRNA	LAMININ ALPHA-1 CHAIN PRECURSOR (LAMININ A CHAIN)	wr65e08.xf NCI_CGAP_Kld11 Hamo septens cDNA clone IMAGE:24845023	Homo sapiens mRNA for KIAA0718 protein, partial cds	EOMESODERMIN	601896233F1 NIH_MGC_19 Homo seplens cDNA clone IMAGE:4125584 6"	Mus musculus potasstum large conductance pH-sensitive channel, subfamily M, alpha member 3 (Konmas), mRNA	Pyrococcus horikoshii OT3 genomio DNA, 644001-777000 nt. postiton (3(7)	H.vulgare Ss1 gene for sucrose synthase	Homo sapiens genomic region containing hypervariable minisatellites chromosome 10[10q28.3] of Homo	saplens	Drosophila melanogaster tiggrin mRNA, complete cds	Homo sapiens microsomal epoxide hydralase (EPHX1) gene, complete cds	602085136F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4248377 5	602085136F1 NIH_MGC_83 Home sapiens cDNA clone IMAGE:4249377 5	Thermologa martima section 85 of 136 of the complete genome	CYSTATHIONINE BETALYASE PRECURSOR (CBL) (BETA-CYSTATHIONASE) (CYSTEINE LYASE)	Homo sapiens mRNA for FLJ00013 protein, partial ods	Homo sapiens mRNA for FLJ00013 protein, partial cds	Homo saplens mRNA for FLJ00013 protein, partial cds	Homo sapians mRNA for FLJ00013 protein, partial cos	X/28407.X1 Sogres_NFL_T_GBC_S1 Hamo sepiens CDNA cione INA/GE:2214.223 3' SETRIER TO SW:C211_HUMAN P63801 PUTATIVE SURFACE GLYCOPROTEIN C210RF1 PRECURSOR;	Homo sapiens hypothetical protein FLJ13220 (FLJ13220), mRNA	yo20e06.r1 Stratagene king (#837210) Hamo sapiens cuna Gone invage: 01.200 o similia to containe MER29 repetitive element	Homo saplens chromosome 21 segment HS21C008	RC3-FN0155-080700-011-d10 FN0155 Homo sapiens cONA	ROS-UM0015-210200-021-A10 UM0015 Hamo saplens cDNA	M.musculus S-antigen gene promoter region	
Top Hit Database Source		ISSPROT	EST_HUMAN		SWISSPROT	EST_HUMAN								EST_HUMAN	EST_HUMAN		SWISSPROT					EST_HUMAN		EST_HUMAN	1	EST HUMAN.	EST_HUMAN	L	SWISSPROT
	토	S	ES	N	8	ES	N	Ę	Z		Z	Z	Ł	S	ES	Ę	જ	¥	Z	Z	노	33	띯	ŭ	Z	Ш	Ü	Ż	S
Top Hit Acession No.	4508862 NT	P10137	Al984806.1	AB018281.1	P79944	BF312963.1	TN11420564	AP00000	10		AL096806.1	U09506.1	AF253417.1		BF678085.1	AE001773.1	P63780	AK024424.1	AK024424.1		AK024424.1			T57160.1	AL163208.2	BE839514.1	AW794952.1	X59799.1	026457
Most Similar (Top) Hit BLAST E Value	3.9E-02	3.7E-02	3.7E-02	3.7E-02	3.7E-02	3.7E-02	3 7E 02	3.6E-02	3.6E-02		3.6E-02	3.5E-02	3.5E-02	3.5E-02	3.5E-02	3.5E-02	3.5E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	3.4E-02	L		Ш
Expression Signal	1.61	6.31	5.07	1.04	1,04	3.76	4	43.88	80		0.73	1.8	1.11	1.01	1.01		1.13							1.86				"	2.79
ORF SEQ ID NO:		11025	12274	12586	13008			13160	13583		13590	L				L	14165					j		42422					
SEQ ID	7692				İ		1			1	8585	1	1	1	1		9182	L				l	L	1	1	1	L		Н
Probe SEQ ID NO:	2632	977	2175	2502	2976	2978	or oc	3115	3870	2	3578	884	8	1531	1531	4092	4189	573	573	574	574	1934	1187	3	32.48	2000	2000	4467	4892

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Single Exon Probes Expressed in HBL100 Cells

		Т	Т	T	Т	7	Т	<b>T</b>	T	Т	Т	Т	Т	T	Ť	Т	Т	$\overline{}$	Т		P	E	Ţ	Ť		15	7	7	4	ť	P	F	H61
	Top Hit Descriptor	Ceanahabditis elegans mRNA for DYS-1 protein partiel	275e08.s1 Sogres testis NHT Homo seriens cDNA clone IMAGE 728408 9	Oricefulus griseus CYP2A17 mRNA for cytochrome P450 2A17 commission cyto	Homo saplens skeletal muscle LIM-protein 1 (FHI 1) nane complete ode	y/25c09.r1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE-127988 F.	Homo saplens skeletal muscle LIM-protein 1 (FHL1) cana. complete cds.	Mus musculus tumor relection antigen ea98 (Tra1), mRNA	Orciolegus cuniculus gene encoding liest sodium-denandent bile edd fransporter	Drosophila melanogastar heat shock protein 68 (han88) gans han884 allala complete ada	Orosophila melanogastar heat shock protein 88 (hsp68) gene, hsp68d allele complete or	LARGE TEGUMENT PROTEIN	601442431F1 NIH_MGC_65 Hamo sepiens aDNA date IMAGE:3848727 5	Homo explens chromosome 21 segment HS21 Cn03	S.cerevisiae chromosome IV reading frame ORF YDL055c	S.cerevisiee chromosome IV reading frame ORF VDI 055c	H. sapiens RP3 gene (XLRP gene 3)	Saviffores rigidifies matures (most) same statements at	Vitreoscilla so, outer membrane andein homolog dans complete cde. The revesees budges maken	partial cds; and unknown genes	Homo seplens duel specificity phosphatase 4 (DUSP4) mRNA	NEURONAL ACETYLCHOLINE RECEPTOR PROTEIN. ALPHA-3 CHAIN PRECLIFSOR (GEAL PHA-3)	Mus musculus adaptor-related protein complex AP-3, delta subunit (Ap3d), mRNA	AU119006 HEMBA1 Homo septens cDNA clone HEMBA1004842 6'	Homo saplens fibrinogen-like 2 (FGL2), mRNA	Pityokteines minutus cytochrome oxidase I gene, partial cds; mitochondrial gene for mitochondrial exonus	z85h03.r1 Soares, testis, NHT Homo sapiens cDNA clone IMAGE:727253 5	Saccharomyces cerevislae stam-loop mutation supressor SSL2 gene, complete ods	Pseudomonas fluorescens family il aminotransferase dene, complete cds	QV2-ST0296-160200-040-e09 ST0298 Hamo sapiens cDNA	Homo sapiens neuropilin 2 (NRP2) gene, complete cds, alternatively spliced	Homo saplens neuropilin 2 (NRP2) gene, complete cds, alternatively spliced	
-1-	Top Hit Database Source	N.	EST HUMAN	LN LN	N I	EST HUMAN	F	LN LN	NT	Į.	TN	SWISSPROT	EST_HUMAN	Z	Z	Z	Z	 		Z	Ę	SWISSPROT	LΝ	EST_HUMAN	ΝΤ	— <b>₽</b>	EST_HUMAN	FX	N.	EST_HUMAN	N	LN	
	Top Hit Acesslon No.	3.4E-02 AJ012469.1					<b>-</b> -	6755862 NT	002005.1	096275.1				_						3.2E-02 AF067083.1	4503416	8845	6871664 NT	118006.1	5730074 NT		-		_			ŀ	
	Most Similer (Top) Hit BLAST E Vatue	3.4E-02	3.3E-02	3.3E-02	3.3E-02	3.3E-02	3.3E-02	3.3E-02	3.2E-02 AJ	3.2E-02 AF	3.2E-02	3.2E-02	3.2E-02	3.2E-02	3.2E-02	3.2E-02	3.2E-02 X94768.1	3.25-02		3.2€-02	3.1E-02	3.1E-02 P1	3.1E-02	3.1E-02 AU	3.1E-02	3.0E-02	3.0E-02	3.0E-02	3.0E-02	3.0E-02	3.0E-02	3.0E-02	
	Expression Signal	1.0	15.62	14.2	. 1.34	1.55	2.72	2.08	2.52	14.14	14.14	8.6	13.22	1.3	0.88	0.88	17.68	3.39		2.67	1.62	1.44	0.95	12.58	5.51	11.29	0.99	0.95	2.92	0.72	7.12	7.12	
	ORF SEQ ID NO:	14856		11185	11656		11656	14318	10214	11145	11146		13091	13644	13861	13862		14595		14848		11328	11929		1		12592	13511	13582		14834	14835	
	Exen SEQ ID NO:	9885	6420	6153	6595	7014	6595	8334	6198	6115	6115	7042	8078	8638	8856	8856	9091	9607		9886	6239	6285	884	9117	10041	6581	7477	8494	8576	8665	9864	8884	
	Probe SEQ ID NO:	4908	371	1149	1599	2031	4053	4343	132	1109	1109	2060	3061	3632	3854	3854	4097	4622		4681	1241	1286	1852	4998	5072	1584	2509	3486	3569	3660	4885	4885	

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Homo septiens mitochondrial glutathione reductase and cytosolic glutathione reductase (GRD1) gene, complete cds. alternatively soliced	AU7e10.r1 Soares fetal liver spleen 1NFLS Homo saniems cDNA clone IMAGE-223130 R	MULTIDRUG RESISTANCE-ASSOCIATED PROTEIN 6 (ABC TRANSPORTER MOAT-C) (PABC11) (SMRP)	S.Wigare benC gene for PEP perhandese	S.Widare penCoene for PEP party values	855509.81 Sogres parathyrnid himmy NHHDA Himm senions ADNA Alone 1980042.91	Homo saplens retinal fascin (FSCN2) gene. even 2	Homo sablens retinal fascin (FSCN2) gene amm 2	Raffus novegicus microtubule-associated protein tau (Mant), mRNA	Human germline T-cell receptor beta chain Dopamine-beta-hydroxylase-like, TRY1, TRY2, TRY3	TCRBV27S1P, TCRBV22S1A2N1T, TCRBV8S1A1T, TCRBV7S1A1N2T, TCRBV5S1A1T, TCRBV13S3,	TCRBV6S7P, TCRBV7S3AZT, TCRBV13S2A1T, TCRBV8S2A2PT, TCRBV7S2A1N4T, TCRBV13S9A3S>	Arabidoosis thallane DNA chromosome 4 contraframent No. 8	W86h12rt Spares multiple sciences 2NNM/SP Home emisse child Alone 1840 E-200402 Et	1988h12.r1 Soares multiple acterosis 2NbHMSP Homo september China Nika DE 280487 87	ye39f04.s1 Soares fetal liver spleen 1NFLS Hamo saplens cDNA clane INAGE-120127 3's similar to contains	Alu repetitive element	Homo sapiens chromosome 21 segment HS21C082	IL3-C70219-280100-062-C09 C70219 Homo sations aDNA	ab02b02.s1 Stratagene fetal retina 937202 Homo saplens cDNA clone IMAGE:839595 3*	Mus musculus histidine rich calcium binding protein (Hrc), mRNA	Mus musculus histidine rich calcium binding protein (Hrc), mRNA	Mus musculus MHC class III region RD gene, partial cols; Bf, C2, G9A, NG22, G9, HSP70, HSP70, HSC704	and smRNP genes, complete cds; G7A gene, partial cds; and unknown genes	Chicken dorsalin-1 mRNA, complete cds	M.musculus DNA for vimentin-binding fragment VimE7	M.musculus DNA for vimentin-binding fragment VimE7	Delnococcus radiodurans R1 section 151 of 229 of the complete chromosome 1	xa52b04.x1 NCI_CGAP_Sar4 Homo sapiens cDNA clone IMAGE:2570383 3' similar to SW:Y089_HUMAN H	Q16041 HYPOTHETICAL PROTEIN KIAA0069;	on28f08.y5 NCL_CGAP_Lu5 Homo sepiens cDNA clone IMAGE:1557827 6*	<b>5.1</b>
Top Hit Database Source	 	EST HUMAN	SWISSPROT	L	Į.	EST HUMAN	Z	Z	F				Ę	EST HUMAN	EST HUMAN		EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	۲٦.	ζŢ	[	¥	NT	NT	NT	NT		EST HUMAN	EST_HUMAN	•
Top Hit Acession No.	228703.1	2805.1		Г	Γ	2.8E-02 AA782516.1		Γ	761			2.7E-02 U66059.1	2		2.7E-02 N47258.1					2.6E-02 AA490021.1	6754241 NT	8754241 NT		109906.1				2.6E-02 AE002014.1		2.6E-02 AW241154.1	٦	
Most Similar (Top) Hit BLAST E Value	2.9E-02	2.9E-02 H7	2.9E-02	2.9E-02	2.9E-02	2.8E-02	2.8E-02	2.8E-02	2.8E-02			2.7E-02	2.7E-02	2.7E-02	2.7E-02		2.7E-02 T95073.1	2.6E-02	2.8E-02	2.6E-02	2.6E-02	2.6E-02		2.6E-02	2.6E-02 L1	2.6E-02	2.6E-02/	2.6E-02		2.6E-02	2.5E-02/	
Expression Signal	1.06	1.37	. 0.73	1.81	1.81	1	1.1	1:1	9.0			1.09	184	2.12	2.12		1.04	0.98	0.83	1.97	1.99	1.89		1.39	3.57	1.27	1.27	1.74		2,61	1.4	
ORF SEQ ID NO:	12459	13836	13881	14812	14813		13323	13324				11613	13374	14060	14081		14980	10598		12388	12400	12401	-		14720	14728	14729	14844		148/4	10566	
Ekan SEĞ ID NO:	7769	8829	8878	9839	6838	7377	8297	8297	9178	*		454	8356	8073	9073		<del>1</del> <del>2</del> <del>2</del> <del>2</del>	929	6347	7280	7282	7282	1000		\$2g	9743	9743	9876	9	830	2003	
Probe SEQ ID NO:	2368	3827	3877	4858	4858	2406	3286	3286	4185			1457	3347	4079	4079		ğ	999	1350	2305	2307	2307	20.45	<b>2</b> 5	*	4759	4769	4897	,	178	828	

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	on26f06.y5 NCI_CGAP_Lu6 Hamo sapiens cDNA clane IMAGE:1657827 6"	601680305R2 NIH_MGC_83 Hamo saplens cDNA clane IMAGE:3950665 3'	601880305R2 NIH_MGC_83 Hamo sapiens cDNA clane IMAGE:3850865 3'	Rattus norvegicus rabphilin-3A mRNA, complete cds	H.carterae mRNA for fucoxanthin chlorophyll a/c binding protein, Fop1	H.carterae mRNA for fucoxanithin chlorophyll alc binding protein, Fop1	PM2-NN0128-080700-001-a12 NN0128 Homo sapiens cDNA	PM2-NN0128-080700-001-a12 NN0128 Homo saplens cDNA	hr3th08.x1 Scares_NFL_T_GBC_S1 Homo expiens cDNA done IMAGE:28340163'	tc72c07.x1 Sogres_NhHMPu_S1 Hamo sepiens cDNA clane IMAGE:20701683'	y 75f11.r1 Sceres fetal liver spleen 1NFLS Homo sapiens cDNA done IMAGE:211149 6	H-2 CLASS I HISTOCOMPATIBILITY ANTIGEN, K-B ALPHA CHAIN PRECURSOR (H-2K(B))	H-2 CLASS I HISTOCOMPATIBILITY ANTIGEN, K-B ALPHA CHAIN PRECURSOR (H-2K(B))	T.thermophila calclum-binding 25 kDa (TCBP 25) protein mRNA, complete cds	H-2 CLASS I HISTOCOMPATIBILITY ANTIGEN, K-B ALPHA CHAIN PRECURSOR (H-2K(B))	H-2 CLASS I HISTOCOMPATIBILITY ANTIGEN, K-B ALPHA CHAIN PRECURSOR (H-2K(B))	za84g08.r1 Soares_fetal_lung_NbHL19W Homo sapiens cDNA clone IMAGE:299294 5	4 Homo sapiens mammary tumor-associated protein INT6 (INT6) gene, exon 4	S.cerevisiae chromosome IV reading frame ORF YDL245c	HSAAACADH P, Human foetal Brain Whole tissue Homo sapiens cDNA	Gallus gallus connexin 45.6 (Cx45.6) gene, complete cds	Gallus gallus connextn 45.6 (Cx45.6) gene, complete cds	CM4-NN0080-290400-160-b04 NN0080 Homo sapiens cDNA	CM3-MT0118-010900-318-g07 MT0118 Homo sapiens cDNA	CM3-MT0118-010900-318-g07 MT0118 Homo sapiens cDNA	xs25d08.x1 NCI_CGAP_Ut2 Home septens cDNA clone IMAGE:2770871 9'	xs25d08.x1 NCI_CGAP_Utz Homo sapiens cDNA clone IMAGE:2770871 3'	601672279F1 NIH_MGC_20 Hamo saplens cDNA clane IMAGE:3955386 5	601672278F1 NIH_MGC_20 Hamo sapiens cDNA clane IMAGE:3955388 5	Homo sapiens KIAA0547 gene product (KIAA0547), mRNA	Rattus norvegicus guanine nucleotide binding protein gamma subunit 11 mRNA, complete cds	Rattus norvegicus guanine nucleotide binding protein gamma subunit 11 mRNA, complete ods	Cotumba livia nucleoside diphosphate kinase (NDPK) gene, nuclear gene encoding mitochondrial protein, complete cds	
AUII FIUDES I	Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	NŢ	NT	NT	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	SWISSPROT	SWISSPROT	NT	SWISSPROT	SWISSPROT	EST_HUMAN	NT.	NT	EST_HUMAN	TN	NT.	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	IN	TN	TN	<b>!</b>	
Single E	Top Hit Acession No.	1783130.1		1.1						2.5E-02 AW592114.1	1378582.1	.1			1		P01901	W05340.1	J94165.1	774293.1	Z20377.1	1.24799.1	124799.1		BE935225.1	BE935225.1	AW 593693.1	AW 583683.1	BF026487.1	BF026487.1		AF257110.1	AF257110.1	AF018287 1	•
	Most Similar (Top) Hit BLAST E Value	2.5E-02 A	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.5E-02	2.4E-02	2.4E-02	2.4E-02 P01901	2.4E-02 P01901	2.4E-02	2.4E-02 P01901	2.4E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.3E-02	2.2E.02	
	Expression Signal	1.4	12.78	4.21	1.83	3.25	3.25	0.85	0.85	5.51	0.68	1.86	1.31	1.31	1.46	1.56	1.56	3.68	5.18	1.88	5.37	0.78	0.76	1.14	0.85	0.85	0.82	0.82	2.76	2.76	0.95	77.0	0.77	20.0	7.90
	ORF SEQ ID NO:	10567	10851	10918		12924	12925	13923	13924	14065	10249	11622	12082	12083	14220	14362	14363			12382	13613	14009		14275	14301	14302	14303	14304	14434			14931	L		
	Exan SEQ ID NO:	5563	5821	5877	7847	7904		10049	10049		5237	0958	7760		1628	9381	9381	6824	6837	7264	8605		L		9319	L	10051		9453	L			L		10/0
	Probe SEQ ID NO:	528	8	858	2689	2885	2885	3932	3932	4083	173	1563	1993	1993	4243	4390	4390	1834	1848	2289	3598	4026	4026	4296	4327	4327	4328	4328	4463	4463	4860	4977	4977	-	97

Page 48 of 209 Table 4 Single Exon Probes Expressed in HBL100

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	Top Hit Descriptor	Homo sapiens chromodoman helicase DNA binding protein 2 (CHD2) mRNA	MYOSIN LIGHT CHAIN KINASE, SKELETAL MUSCLE (MLCK)	MYOSIN LIGHT CHAIN KINASE, SKELETAL MUSCLE (MLCK)	S nneumoniae pood gene and open reading frames	Chinamical No. 1 No. 1 No. 1 Crap Gast Home saciens cond clone IMAGE:1084782 3	Internative hereal disease wine segment Batrein II 4 VP1 cans. complete cds	INTEGRALS DATE OF THE SERVICE CONTRACT OF THE SERVICE	Principal Control Control of Cont	S.CORONSIBO GUICHICON IN COMMING HOME CAN ADDRESSOR	AV761502 MILO FIGURE COLON GUITA MILONICO I O	Dictyostelium discoldeum nistignine kinese C (dinko) minara, unimate cus	Bacillus subtilis cotKLM cluster, CotK (cotK), CotL (cotL), and space contiguous cotKLM cluster, CotK (cotK), CotL	COMPAND UND SHIED DE MATERY DEOTEIN ROA	KERALIN, HIGH-SULTUR MALITAN FROITEIN, DZA	KEKALIN, HIGH-SULTUR MATRIX PROTEIN, BZA	KERATIN, HIGH-SOLL-UK MATRIX-TOOLEIN, BZA	JAKSHU7,71 Soares melanocyle ziviznim ricinio septens control	2030UST Source IV modified frame OPF VDI 2450	S.Geenstee Gridingsonte IV Teatury Hans Son 15-170	OUZUISSUOTING COOK DIMOTING CONTROL CO	Borrella burgdorten plasmid choz-z, etc. etc. etc. etc. etc. etc. etc. etc.	Wgo Idi I.M. Sowies Andrews Andrews for heir keretin, extent 2 to 7	Tomo sapidie pusare pain in possessing a	Autistical at Constant Wilms hims Home actions CDNA clone IMAGE:1126918 3'	TELEGO CANCIL COAD DAY Home series CONA close IMAGE:3309998 3' similar to contains MER1.13	rganwa.r i norgona ji norgona ji norgona ka	QV4-NN0038-270400-187-h05 NN0038 Homo septens CUNA	Mus musculus DinB homolog 1 (E. coll) (Dinb1), mixNA	Ba15010.11 Coares, Ninthing of Thomas express of the Coares and the Coares of the Coar	Mus musculus Ulus fromadog 1 (E. cuil (Ulim), ilinus	Home sapiens genomic region containing hypervariable minisationies circlinocure. It proceds to the containing t	Sapkens Home caniers twodhedical protein FLJ10379 (FLJ10379), mRNA		•
	Top Hit Database Source	5	SWISSPROT	SWISSPROT	III	TANK IN TOTAL	NOWAL TOWAR	IN	EST_HUMAN	□ <b>L</b> N	EST HUMAN	K		IN.	SWISSPROT	SWISSPROT	SWISSPROT	EST HUMAN	EST HOMAN	Z	EST HOMAN	NT	EST HUMAN	Z	Į.	ESI HOMAN	EST_HUMAN	EST_HUMAN	M	EST HUMAN	NT	!	<u> </u>	Z	<u></u>
	Top Hit Acession No.	4557448 NT	7244	7343		1	2.2E-02 AA5/7/85.1	7	٦	2.2E-02 Z74283.1				_			P02438	N29266.1	2.1E-02 AA481271.1	2.1E-02 Z74293.1	2.1E-02 BF343655.1	U44914.1				AA665737.1	BF002932.1	12		AA45653	6753635 NT		AL 08680		1 N 1 1 8 2 2 3 9 1 N 1
- FILE CO.	Most Similar (Top) Hit 1 BLAST E Value	2.2E-02	00 00 00 0	2 2E 02 Bu		2.2E-02 282001.1	2.2E-02 /	2.2E-02/	2.2E-02 /	2.2E-02	2.1E-02/	2.1E-02/		2.1E-02	2.1E-02 P02438	2.1E-02 P02438	2.1E-02 P02438	2.1E-02 N29266.1	2.1E-02	2.1E-02	2.1E-02	2.1E-02 U	2.1E-02	2.1E-02	2.1E-02	2.1E-02	2 0E-02								2.0E-02
	Expression Signal	12	100	0.00	6.97	1.43	1.88	3.57	1.18	0.68	4.33	7.7		7.23	1.17	1.17	1.17	3.7	0.94	0.67	9.0	1.92	1.35	0.68	5.38	0.76	4.78	8.6							1.31
	ORF SEQ ID NO:	1	1	98/1	11/80	12053			13769	13829				11283	11816	11817	11818	10820	13524	13993	14159	14297	14310		14549	14569	40000						11110	11219	11220
	SEQ ID	8703		6718	81/9	6950	8359	8998	8766		5453			6242		L	L			L	9175	L	8324	9328	9560	9229		2008	Ŀ	L		L	1 6079	0 6182	0 6182
	SEQ ID	4700	2	1723	1723	1965	3350	3561	3763	3820	416	448		1244	1744	1744	1744	2744	3502	4009	4182	4322	4333	4540	4572	4591	'	2   \$	257	i k			1071	1180	1180

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Single Exon Probes Expressed in No. Cells	Top Hit Descriptor	Homo sapiens hypothetical protein FLJ10488 (FLJ10488), mRNA	Homo saplens hypothetical protein FLJ10486 (FLJ10486), mRNA	Arabidopsis thallana DNA chromosome 4, contig fragment No. 32	7g51c08.x1 NCI_CGAP_Pr28 Homo sepiens cDNA clone IMAGE:3309998 3' straiter to contains MER1.t3 MER1 repetitive element ;	Mus musculus sema domain, transmembrane domain (TM), and cyloplasmic domain, (semaphorin) 6B	(Semaol), mysky	Arabidopsis mailaria CZTZ zno mosa promini ZZ michak companio cus	P. vulgaris hydroxyproline-rich glycoprotein (HRGP) mRNA, 3' end	lg183e03.x1 NCI_CGAP_Kid3 Homo sapiens cDNA clone IMAGE:1886076 3"	inf19a07.s1 NCI_CGAP_Prf Homo sepiens cDNA clone IMAGE:914198 similar to contains L1.f1 L1	Home eaclers chromosome 21 segment MS21C103	Library continue chromosome 24 soumed HS4C403		Arabidopsis mailana Lina chromosome 4, congrir ingmentino, bu	nw04f05.s1 NCI_CGAP_SS1 Homo saplens cLINA clone IMAGE:1238337 3	AV648669 GLC Homo sapiens cDNA clone GLCBLH07 3	Urotrichus talpoides mitochondrial gene for cytochrome b, complete cds	yz28b02.s1 Soares_multiple_sclerosis_2NbHMSP Homo sepiens cDNA done IMAGE:284331 3*	601572682F1 NIH_MGC_67 Homo saplens cDNA clone IMAGE:3839564 5	qn04c07.x1 NCL_CGAP_Lu5 Homo sapiens cDNA clone IMAGE:1897260 3' straitar to contains Alu repetitive	(Mucroscome Imitane VIHA1 measuress (vIHA1) and VIHA2 presuress (vIHA2) comes. Dertiel cds	Injurication became with the second s		1	DNA clone IMAGE:2144661 3' similar to	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 50	hn52c08x1 NCI_CGAP_Co17 Homo sapiens cDNA clone IMAGE:3027274 3' similar to contains element	MER29 repetitive element;	H.francisci mRNA for myelin basic protein (MBP)		te52x09.x1 Scares_NFL_T_GBC_S1 Hamo saplens aDNA clone IMAGE::2090/296 3		
ON PRODES D	Top Hit Database Source	<b>_</b>	1	1	EST_HUMAN	 			뉟	EST_HUMAN	CT LUMBAN	PANOL 121		Į.	Ę	EST HUMAN	ST HUMAN	Ę	EST HUMAN	EST_HUMAN	- For		I N	SWISSPROI	SWISSPROT	EST_HUMAN	Į.	_	EST HUMAN	NT	IN	EST_HUMAN	EST HUMAN	
	Top Hit Acession No.	8922453 NT	8922463 NT	4L161632.2 NT	002832.1		4,4	-		2.0E-02 AI271995.1		1.9E-02 rw3/2/04:1	I	1.9E-02 AL163303.2	١	1.9E-02 AA713856.1 E						T	1.9E-02 AF141940.1	٦			1 9E-02 AL 181550.2					1.8E-02 AI805829.1	Г	7
	Most Similar (Top) Hit BLAST E Value	2.0E-02	2.0E-02	2.0E-02 AL	2.0E-02		2.05-02	2.0E-02	2.0E-02	2.0E-02	4 00 00	1.85-02	1.95-02	1.95-02	1.96-02	1.9E-02	1.9E-02	1.9E-02	1.9E-02	1.9E-02		ı	ļ		1.9E-02	1.95-02				L				12
3	Expression Signal	1.69	1.69	1.8	1.66		221	4.5	1.61	0.84	97.	1./0	2.10	2.16	1.18	7.79	1.67	0.75	0.91	8.78			1.5	1.75	1.75	281			1.36	1.42				
	ORF SEQ ID NO:	11912	11913		10080				13898							12874	12922								14054	14385	۱		10403		L		13804	
	SEO ID NO:	9859	6825	7683	5097		8084	8172	8900	9920		90/6	29/3	١	ı	7855	7900	8201	8536	8629					9064	8	1	١	5396		L		1	
	Probe SEO ID NO:	1835	1835	2728	3005		3874	3156	3900	4943		ğ	<u>88</u>	88	2434	2835	2881	3185	3530	3622		3633	3934	4070	4070	4410	200	7004	4	1165	2807	3438	200	3/93

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	MR1-0T0011-280300-009-g04 OT0011 Homo sepiens cDNA	ak24h04.s1 Soares_bests_NHT Homo saplens cDNA clone IMAGE:1406835 3'	QV4-DT0021-301289-071-b11 DT0021 Homo sepiens cDNA	HYPOTHETICAL PROTEIN DJ845024.2	601310626F1 NIH_MGC_44 Homo septens cDNA clone IMAGE:3632160 5	h734s03.x1 Soares_NFL_T_GBC_S1 Homo saptens cDNA clone IMAGE::2833740 3' similar to contains L1.t1 L1 repetitive element;	h734803.x1 Soares_NFL_T_GBC_S1 Homo explens cDNA clone IMAGE:2833740 3' similar to contains	L1.t1 L1 repeditive element;	Homo sapiens chromosome 21 segment HS21C004	Oryctolagus cuntculus mRNA for mitsugianin28, complete cds	Homo sapiens putative Rab5 GDP/GTP exchange factor homotogue (RABEXS), mRNA	qb22a08.x1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:1696982 3*	hm45e04.x1 NCI_CGAP_RDF1 Homo septens cDNA clone IMAGE:3015534 3' similar to contains	MER19.b1 MER19 repetitive element;	ac19f04.s1 Stratagene ovary (#837217) Homo sapiens cDNA clone IMAGE:856927 3' similar to contains Alu repetitive element,contains element MER24 repetitive element ;	ye86f08.r1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:124647 6	qm08g07x1 NCI_CGAP_Lu5 Homo sepiens cDNA clone IMAGE:1881276 3' similar to gb;X62359 ZINC FINGER PROTEIN 30 (HUMAN);	hi34a03.x1 Soares_NFL_T_GBC_S1 Homo septens cDNA clone IMAGE:2833740 3' strutter to contains	L1.f1 L1 repetitive element;	Messenger RNA for anglerfish (Lophius americanus) somatostatin li	ov51e02.s1 Soares_testis_NHT Homo septens cDNA clone IMAGE:16408583	Mycobacterium tuberculosis H37Rv complete genome; segment 13/162	Treponema maltophilum flaB2, flaB3 and filD genes for flagellin subunit proteins and CAP protein homologue	LIVER CARBOXYLESTERASE 22 PRECURSOR (EGASYN) (ESTERASE-22)	LIVER CARBOXYLESTERASE 22 PRECURSOR (EGASW) (ESTERASE-22)	ne81d06.s1 NCI_CGAP_Ew1 Homo saplens cDNA clone IMAGE:910867	Homo sepiens mRNA for KIAA0634 protein, partial cds	s; mitochondrial gene for mitochondrial product	IL3-CT0218-160200-063-C07 CT0219 Hamo saplens cDNA	
2000	Top Hit Database Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	SWISSPROT	EST_HUMAN	EST HUMAN		EST HUMAN				EST_HUMAN		EST HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN		EST HUMAN		EST HUMAN			SWISSPROT	SWISSPROT	EST HUMAN			EST HUMAN	
3		ES	ES	ES	MS	ES.	ES.		ES	NT	NT	NT	ES.		ES	ES	ES	S		ES	Z	ES	Ž	눌	8	S	123	볼	눌	Sil.	
DIRING.	Top Hit Acession No.	1.8E-02 AW879122.1	1.8E-02 AA861446.1	1.8E-02 AW936363.1	060810	1.7E-02 BE394869.1	1.7E-02 AW573183.1		1.7E-02 AW573183.1	1.7E-02 AL163204.2	1.7E-02 AB004816.1	7857495 NT	1147615		AW827368.1	AA669618.1	1.7E-02 R02506.1	Al305279.1			V00641.1	AI015076.1	AL021929.1	Y18889.1	'n					AW850652.1	
	Most Similar (Top) Hit BLAST E Value	1.8E-02	1.8E-02	1.8E-02	1.8E-02 O60810	1.7E-02	1.7E-02		1.7E-02	1.7E-02	1.7E-02	1.7E-02	1.7E-02 A		1.7E-02 A	1.7E-02	1.7E-02	1.7E-02 A		1.7E-02	1.7E-02	1.7E-02	1.6E-02	1.65-02	1.6E-02	1.6E-02	1.6E-02	1 6F-02	1.6E-02	1.6E-02	
	Expression Signal	16.0	1.08	1.59	96.0	1.21	2.15		2.15	2.15	7.25	1.47	1.09		4.78	1.04	1.86	1.34		1.47	1.82	6.38	1.83	1 13	1.01	1.01	1.12	0.95	0.73	5.11	
	ORF SEQ ID NO:	13802		14284	14765	10952	11828		11829				12966					14300		14370	14552			14680	L	L			12987		
	Exan SEQ ID NO:	8796	6968	9298	9782	5913	6747	L	6747	6823		7532		١.	<b>8</b>	9041	L	ļ		9387	L		١.		L					L	
	Probe SEQ ID NO:	3783	3970	4306	4798	895	1753		1763	1833	2054	2569	2929		3433	4045	4077	433.2		4396	4576	4668	507	484	2487	2487	2577	i S	205	\$	

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Hamo sepiens chromosome 21 segment HS21C101	Mus musculus major histocompatibility complex region NGZ7, NG28, RPS28, NADH oxidoreductase, NG29, KIFC1, Fas-binding protein, BING1, tapasin, RafGDS-like, KE2, BING4, beta 1,3-galactosyl transferase, and RPS18 canes. complete clast Second name, nartials.	Homo sapiens transcription factor (HSA130894), mRNA	W27b07.s1 Soeres (et al liver soleen 1NFLS Homo sentena cityna cityna itiAACE-343coa 9	Arabidopsis thatiana DNA chromosome 4. conting fragment No. 20	Homo sepiens CACNA1F gene, exons 1 to 48	Homo sapiens CACNA1F gene, exons 1 to 48	MR4-TN0115-080900-201-b12 TN0116 Homo seplens cDNA	Homo saplens eukaryotic translation initiation factor 4E (EIE4E) mRNA	Chlamydophila pneumoniae AR39, section 58 of 84 of the complete genome	Homo sapiens NESH protein (LOC51225), mRNA	Haemophilus Influenzae Rd section 115 of 163 of the complete genome	Xenopus laevis neurodenin related 1b (X-NGNR-1b) mRNA commisse and	AV723785 HTB Homo septens cDNA clone HTBAHH11 5	Homo sepiens down-regulated in metastasis (DRIM), mRNA	Bifidobacterium ka+/IH+ entiporter (nhaB), cytosine deaminese, and alpha-galactosidese (egil.)	genes, complete cust, and N-acety/glucosamine/xylose repressor protein (nagC/xylR) gene, partial cust	Arehdonele thellene DNA chromosome 4 contra forces and one	Arabidoosis thaliana DNA chromosome 4, contig fragulatin No. 62	Mus musculus histocompatibility 2, complement component factor B (H2-Bf), mRNA	EST374761 MAGE resequences, MAGG Hamo sapiens cDNA	EST374761 MAGE resequences, MAGG Homo saplens cDNA	601567403F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3842280 6"	601667403F1 NIH_MGC_21 Hame saplens cDNA clone IMAGE:3842280 6'	601558462F1 NIH_MGC_58 Homo saplens cDNA clone IMAGE:3826335 6	Homo sapiens chromosome 21 segment HS21C001	Oenothera berterlana NADH dehydrogenase subunit 2 (nad2) gene, exons 1-2	602129475F1 NIH_MGC_56 Hamo saplens cDNA clone IMAGE:4288203 6		Mus musculus beta-sarcoglycan gene, complete cds	
	Top Hit Databese Source	M	<u> </u>	5	EST HUMAN	F	Į.	¥	EST_HUMAN	1.	¥	1	F	LN	EST HUMAN	1		N.FOT LIMAN	LA LA	Ę	П	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	NT	EST_HUMAN	EST HUMAN	NT	
	Top Hit Acession No.	.163301.2	1.6E-02 AF110520.1	1923734	1.5E-02 N39521.1	161694.2	006216.1	106216.1	1.5E-02 BF092942.1	4503534 NT	1.4E-02 AE002230.2	TN05980 NT			1.4E-02 AV723785.1	37040		T	Τ	1.4E-02 AL161586.2	TN 8169669											·
	Most Similar (Top) Hit BLAST E Value	1.6E-02 AL	1.6E-02	1.5E-02	1.5E-02	1.6E-02	1.5E-02 AJC	1.5E-02	1.5E-02	1.5E-02	1.4E-02	1.4E-02	1.4E-02 U32800.1	1.4E-02	1.4E-02	1.4E-02	Ļ	1.4C-02/	1 4F-02	1.4E-02/	1.4E-02	1.4E-02	1.4E-02 /	1.4E-02	1.4E-02	1.3E-02	1.3E-02	1.3E-02	1.3E-02	1.3E-02	1.3E-02	
	Expression Signal	1.1	1.82	23.98	2.42	1.29	1.24	1.24	0.88	0.97	1.25	3.55	1.37	2.38	96'0	0.87		7 2 2	6.12	6.12	10.27	7.08	7.06	6.63	6.63	1.4	1.7	0.87	1.9	1.9	1.18	
	ORF SEQ ID NO:	13761			12171			13018	13654	14952		11138				12962	455	13347	13439	13440	13585	14331	14332	14695	14696		11894	12970	13178	13179		
	Exon SEQ ID NO:	8761	9047				8005	8005	8647	220	5451	6108	6235	6278	6483	7945	93.0	8325	8413	8413	8228	9352	8352	9711	9711	6818	6901	7953	8157	8157	8864	
	Probe SEQ ID NO:	3758	4051	742	2081	2110	2987	2987	3641	5006	414	1101	1237	1279	1486	2926	6	3314	3404	3404	3572	4361	4361	4726	4726	1828	1915	2934	3141	3141	3862	

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Table 4
Single Exon Probes Expressed in HBL100 Cells

					_		_	_		_	_		_	2	<u> </u>	Ļ	4	4	S		D;	1	4	H	H	ĘE
Top Hit Descriptor	Human germiina T-ceil receptor beta chein TCRBV17S1A1T, TCRBV2S1, TCRBV10S1P, TCRBV29S1P, TCRBV19S1P, TCRBV19S1P, TCRBV19S1P, TCRBV19S1P, TCRBV19S1, TCRBV14S1, TCRBV14S1, TCRBV4S1A1T, TRY4, TRY5, TRY6, TRY7, TRY8, TCRBV1S1, TCRBJ1S1, TCRBJ1S2,>	265g01.r1 Soares retina N2b4HR Homo sapiens cDNA clone IMAGE:381840 6' similar to contains element L1 repetitive element ;	HYPOTHETICAL 17.1 KD PROTEIN IN PUR5 3'REGION	qd68e12.x1 Sogres_testis_NHT Hamo sepiens cDNA clone IMAGE:1734670 S' similer to contaîns L1.f1 L1 repetitive element ;	Homo sapiens chromosome 21 segment HS21C013	AV731704 HTF Hamo saplens cDNA alone HTFBHG11 5	x/37e09.x1 Soares_NFL_T_GBC_S1 Hamp sepiens cDNA clane IMAGE:2659432 3'	x/37e09.x1 Scares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2659432 3'	zm88e03.r1 Stratagene ovarian cencer (#337219) Homo sapiens cDNA clone IMAGE:645020 6	y11b08.s1 Soares placenta Nb2HP Homo saplens cDNA clone IMAGE:138903 3'	Mus musculus interferon regulatory factor 5 (Irf5), mRNA	Human hereditary haemochromatosts region, histone 2A-like protein gene, hereditary haemochromatosts (HLA-H) gene, RoRet gene, and sodium phosphata transporter (NPT3) gene, complete cds	Cynops pyrrhogaster CpUbiqT mRNA, partial cds	AV731704 HTF Homo sapiens cDNA clone HTFBHG11 5'	Mus musculus POZ/zinc finger transcription factor ODA-8 mRNA, complete ods	zm69e11.s1 Stratagene neurocpithelium (#937231) Homo sapiens cDNA clone IMAGE:630924 3	H.sapiens LIPA gene, exon 4	H.sapiens LIPA gene, exon 4	602018037F1 NCI_CGAP_Bm87 Homo septens cDNA clone IMAGE:4153808 6	za40e05.r1 Soares fetal liver spiean 1NFLS Homo sapiens cONA clone IMAGE:286040 5	In96510.x1 NCI_CGAP_Ov23 Hamo sepiens cDNA clane IMAGE:2216639 3' similer to SW:XPF_HUMAN Q92889 DNA-REPAIR PROTEIN COMPLEMENTING XP-F CELL;	RC3-ST0197-120200-015-g11 ST0197 Homo saplens cDNA	DKFZp586E0924_s1 586 (synonym: hute1) Homo sapiens cDNA clone DKFZp588E0924	MR3-CT0176-111099-003-e10 CT0176 Homo saplens cDNA	oc22h08.s1 NCI_CGAP_GCB1 Homo septens cDNA clone IMAGE:1350495 3'	RC0-FN0025-250500-021-d02 FN0025 Homo saplens CDNA
Top Hit Database Source	<u> </u>	EST_HUMAN	SWISSPROT	EST_HUMAN	M	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	LNT L	¥	Į.	EST_HUMAN	NT	EST HUMAN	·	NT	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN
Top Hit Acession No.	1.3E-02 U88081.1 N	100		2.1		1.2E-02 AV731704.1 E					4367	1.2E-02 U91328.1	-	Г				П		1.1E-02 N99523.1	1.1E-02 AI663508.1	_				
Most Similar (Top) Hit BLAST E Value	1.3E-02	1.2E-02	1.2E-02 P38898	1.2E-02	1.2E-02	1.2E-02	1.2E-02	1.2E-02	1.2E-02	1.2E-02	1.2E-02	1.2E-02	1.2E-02	1.2E-02	1.2E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.1E-02	1.0E-02		
Expression Signal	1.07	4.04	1.62	6.89	127	1.07	1.15	1.16	6.83	1.97	0.95	. 3.86	1.51	2.45	1.29	1.26	1.43	1.43	3.37	4.07	3.11	0.69	2.3	8.28	0.95	2.95
ORF SEQ ID NO:	14742	10414		10773	12208		12468			13251	14699	14731		14865		11289	11739	11740			13478		14650			13044
Exan SEQ ID NO:	9755	5404	5487	5752	7092	7095	7347	7347	8045	8230	9714	9745	9855	9891	9666	6248	6684	8684	6972	7828	8452	8988	8996	5087		Ш
SEQ ID	1,5	38	₹ 18	222	2112	2116	2375	2564	3028	3215	4729	4761	4876	4912	5025	1250	1688	1668	1987	2808	3444	3990	4683	-	2497	3016

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Table 4
Single Exon Probes Expressed in HBL100 Cells

														_		_		_	_	_	٠,	-4	_	Γ,	4		5	-	<del>#</del>	-	8	$\mathbf{D}$	اد	Ξ.
	Top Hit Descriptor	601648867R1 NIH MGC_74 Hamp sepiens cONA clane IMAGE:3833689 3'	Mus muscalus corticohootin releasing hormone receptor 2 (Orthr2), mRNA	Tring 4 Secretarillians ANEL Sections of the MARE 19983 5	Apphilia Source form investigation appears for the second	Human glycoprotein hormone alpha-subunit (GCA) gene, o namk	wh42/09 x1 NCI_CGAP_Kid11 Home sapiens cDNA clone IMAGE:2383433 3' similar to contains seamon.	MER22 MER22 repetitive element;	601470242F1 NIH MGC 67 Homo septems curva cione invasication cu	Arabidopsis thaliana DNA chromosome 4, contig tragment no. 39	1244e10.y1 NCL CGAP BITSZ Hamo septens CUINA CIONE INVIGE 2231400 0	Mus musculus corticotropin releasing normans receptor 2 (Critiz), minura	2h30e03.s1 Scares_pineal_gland_N3HPG Homo septens cunva cione imagin:413590 5 serius in contente Alu repetitive element;	Homo sapiens adenylosuccinate lyase gene, complete cds	Homo saplens chromosome 21 segment HS21C083	Homo sapiens SCL gane locus	HYPOTHETICAL 127.0 KD PROTEIN IN RAD24-BMH1 INTERGENIC REGION	HYPOTHETICAL 127.0 KD PROTEIN IN RAD24-BMH1 INTERGENIC REGION	QV0-FN0181-140700-304-g10 FN0181 Homo sapiens cDNA	CM4-NN0118-300600-223-b05 NN0119 Homo saplens cDNA	Cryptosporialum pervum HC-10 gene, complete cds	Cryotosporidium parvum HC-10 gene, complete cds	Glycine max glutathione S-transferase GST 21 mRNA, partial cds	AV731712 HTF Homo sepiens cDNA clone HTFAZF10 6'	ab79b09.s1 Stratagene fetat retina 937202 Homo sapiens cDNA clone IMAGE:863145 3	xx21b02.x1 Sogres_NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE:2613/39 3	HISTIDINE-RICH GLYCOPROTEIN PRECURSOR	UI-H-BI3-akb-c-10-0-UI.s1 NCI_CGAP_Sub5 Homo sapiens cDNA clone IMAGE:2/33081 3	Rattus norvegicus neuronal nicctinic acetylcholine receptor subunit (Alpha10) mRNA, complete ods	The state of the s	Dictyosinglism disconded in Indian sealors and Above IMAGE-2809033 3' similar to TR:012987 Q12987	X634/UBX NOL_COM_Off Halls adjusts Colon C	hh89a05,y1 NCI_CGAP_GU1 Homo saptens cDNA clone IMAGE:2889938 o	
	Top Hit Database Source	EST HUMAN			EST HUMAN			EST HUMAN	EST HUMAN		HUMAN		EST HUMAN				SWISSPROT	SWISSPROT	EST HUMAN	EST HUMAN				EST HUMAN	EST HUMAN	EST HUMAN	SWISSPROT	EST HUMAN				EST_HUMAN	EST HUMAN	
Ţ.		EST	5		2	Ę		<u>图</u>	S	토	EST	뉟	· ES	Ę	뉟	뉟	8	8	S	S	Ę	į	E	123	S S	ES	S	12	눌		ᅬ	<u>ಟ</u>	Ш	ľ
	Top Hit Acession No.	BE98899.1	TM MORGATA	0100061	<b>138587.1</b>	.05832.1		9.0E-03 AI798128.1	3E781889.1	L161559.2	3E047949.1	6753521 NT	8.0E-03 AA723007.1	8 0E-03 AF-108858.1	8 0F-03 AL 163283.2	8.0E-03 AJ131016.1	232844	232644	8 0F-03 BF840049.1	A 0E-03 RF3R3327 1	AE097183.1	A E007483 4	7.0E-03 AF243376.1	AV731712.1		AW303599.1	P04929	AW44463.1	AF196344.1		U60086.1	AW117711.1		
	Most Similar (Top) Hit BLAST E Value	1.0E-02	20.	1.UE-02	1.0E-02 R98567.1	1.0E-02 LO		9.0E-03	9.0E-03	9.0E-03	8.0E-03	9.0E-03	8.0E-03	8 0F-03	8 OF -03	8.0E-03	8 OF-03	8.0E-03 P32644	8 0F-03	B OF OS	7 OF 03 A	7 05 00 7	7 OF 03	7.0E-03	7.0E-03	7.0E-03	7.0E-03	7.0E-03	<u> </u>		7.0E-03	7.0E-03		ĺ
	Expression Signal	88		8.4	3.4	1.72		2.15	1.46	1.79	1.02	96.0	2.78	52 19	1 30	890	1 23	123	0 05	20.2	44.47	44.47	11.47				1.36	0 0			1.05	1.12	1.1	
	ORF SEQ ID NO:	13228		14608	14665	14810		10941		12425		14970		44023				Ŀ					10/27											
	Exan SEQ ID NO:	8.20A	ı	1	7898	9836		2800	6243			2668		1		L	L	L	1					2007				L			8208		1	
	Probe SEQ ID NO:	9480	200	4632	4697	4855		882	1245	2331	4828	5028	400	22.0	1000	CROS	2020	3504		4134	4238	88	88	708	1088	13/6	14/5	0000	of S	3/22	4213	1	4404	\$

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Hamo explens chromosome 21 segment HS21Cn78	hd22a05.x1 Soares JNF_T_GBC_S1 Homo explens cDNA clane IMAGE:2910224.3' similar to SW-PXR_HUMAN 075489 ORPHAN NICS FAR RECEPTOD BXB.	hd22e05.X1 Soares_NRL_T_GBC_S1 Home septems cDNA done IMAGE:2910224.3' similar to SW:PXR_HUMAN OYFIRD ORDHAN NI ICH EAD DESCENTION DAYS	Danio reno odorani recentor cene chietar	B178e11.s1 Source that NHT Home conjunction about the constant of	87/86/1.s1 Spares lessis NHT Homo seniers child characters of	y77/h04.r1 Sogres fetal liver spleen 1NFLS Home saplens cDNA clane IMAGE 241381 R	Notoncus sp. cytochrome o addess subunit II gene perfiel cyte: mitrobouddol sees for with the contract of the	Fugu rubripes zino finger protein, isotocin, fatty acid binding protein, sepiapterin reductases and vasotocin genes, complete cds	Fugu rubripes zinc finger protein, isotocin, fatty acid binding protein, sepiapterin reductasse and vasotocin	zci3a11 fl Sogres negatived time MUDA U	III.H.Bit.com.c.08.0.11 et NO 100 Bits II.	RC1-BT 1606-2604010 014-607 BT 1608 U.S. September CDNA clone IMAGE:3087764 \$	Mus musculus alucosamina Cabacabate dominana (Cara) - Data	800942904F1 NIH MGC 15 Home seniers CDNA close IMAGE Server	W62h10.s1 Soares multiple sclerosis 2NbHMSP Homo seniens cDNA close 144625-378476 st	043311x1 Scares testis NHT Hamo sapiens CDNA clans IMAGE-16301243	EST27116 Cerebellum II Homo saplens cDNA 6' end similar to EST containing Atu reneet	Human germline UBE1L gene similar to the gene for ublaultin-activating enzyme, exons 1-22	Chlamydia trachomatis pertial ORFB; aminoacyl-tRNA synthase, complete cds; complete ORFA, and grpE- like protein, complete cds	Chlamydia trachomatis partial ORFB; aminoacyl-tRNA synthase, complete cds; complete ORFA, and grpE- like protein, complete cds	Chlamydia trachomatis partial ORFB; aminoacy-tRNA synthase, complete cds; complete ORFA, and grpE- like protein, complete cds	Chlamydia trachomatis partial ORFB; aminoscyl-tRNA synthase, complete cds; complete ORFA, and grpE- life protein, complete cds	Arabidopsis thaliana mRNA for DEAD box RNA halicasa RH3	Homo sepiens mRNA for KIAA1180 protein, partial cds	
2000111101	Top Hit Database Source	N F	EST HUMAN	EST HUMAN	1	EST HUMAN	EST HUMAN	EST_HUMAN				EST HIMAN	EST HIMAN	EST HIMAN		EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN								
1		Z	<u> </u>	l ü	E	Ü	i iii	Ü	<u> </u>	볼	<u> </u>			i jü			I W	I S	ES	INT	<u> </u>	Ż	호	Þ	Z	Ž	
	Top Hit Acession No.	AL 163278.2	AW611148.1	4W611148.1	6.0E-03 AF112374.1	A759135.1	4A759135.1	6.0E-03 H75690.1	6.0E-03 AF190338.1	J90880.1	J80880 4	6.0E-03 W37985.1	6.0E-03 BF510988 4	8.0E-03 BE077358.1	6754029	6.0E-03 BE250108.1	158946.1	6.0E-03 AI016833.1	6.0E-03 AA324242.1	34170.1	25105.1	25106.1	25106.1	25105.1	5.0E-03 AJ010457.1	5.0E-03 AB033006.1	
	Most Similar (Top) Hit BLAST E Value	7.0E-03 AL	6.0E-03 AV	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03	6.0E-03 U90880.1	6.0E-03 U90880 4	6.0E-03	8.0E-03	8.0E-03	6.0E-03	6.0E-03	6.0E-03 N58946.1	6.0E-03	6.0E-03	6.0E-03	5.0E-03  1.25105.1	5.0E-03 L25106.1	6.0E-03   L25106.1	6.0E-03	5.0E-03	5.0E-03	
	Expression Signal	2.67	9.46	. 9.46	1.02	4.19	4.19	2.51	0.85	1.31	1.31	1.13	6.07	<u>1</u>	1.18	98'0	1.31	1.27	6.94	0.98	1.81	1.81	22	22	1.24	2.3	
	ORF SEQ ID NO:		11283	11264	12770	12857	12858			13335	13336		13588	13628	13696				14529		10696	10697	10696	10697	11132	12686	
	Exan SEQ ID NO:	9814	6221	6221	7856	7841	7841	8189	8247	8309	8309	8478	8583	8618	8694	8852	9192	9233	9544	9927	5688	5688	5688	6688	6102	7572	
		4830	1221	1221	2699	2820	2820	3173	3232	3298	3298	3470	3576	3611	3690	3850	4189	4239	4556	4950	681	661	299	882	1095	2810	

Page 55 of 209 Table 4 n Probes Expressed in HBI 10

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Single Exon Frobes Expressed in HBL100 Cells	Top Hit Descriptor		COTTO TANKE TO THE MINISTER OF THE PROPERTY OF	Arabidonele the local DNA character and the company of the company	Alabados u validada DIVA Gruonosome 4, conto fragment No. 3	ylonguz.si Soeres breast ZNBHBst Homo saplens cDNA clone IMAGE:155688 3' Homo saniens nertial IMD states for IMA	Pseudomonas genularings arrein DA.	Citrus sinensis seed chrone provide a letter — DNA	EST12248 (Batis himse) Home sections and an extension of the section of the secti	Citus shensis seed stress awkin eliti — DNA	Homo sablens SCL demo hours	And State of Married Lives and Lives	LILHE-BIND AND AND AND AND AND AND AND ASSESSED SEQUENCES OF A CONTROL OF SECURE OF SE	WELGA Some infection 1 Min Miles. With the septent CDNA clone IMAGE:3076831 6"	PHOSPHATION INDITION SAMES A 2012 MILES OF COMPANY OF THE SAME SAME SAME SAME SAME SAME SAME SAM	075012 \$1 Sours NEI T GRO SALDON STATE THE STATE OF THE S	vo51604 st Spares infant brain 1NIB Uman and an annual and an an an an an an an an an an an an an	73 - 10 101 COR SE IMENINAMI INID FIGURE SEPRENS CLINA CIONE IMAGE: 35988 3'	281808 f1 Strategare Color (#037204) Long explicit Color	RC8-UM0014-170400-073-G01   Monta Home series CONA Good IMAGE: 510888 5	2559a01.11 NCI CGAP GCB1 Home seniors chiving like object to the	AV708305 ADC Homo sapiens cDNA clone ADCAKROR 5'	Safftis nonwerigies hans 4 sethands and effection in the	78140R of Strangers Aloc (4037004) Unactual year in the second of the se	801304161F1 NIH MGC 21 Home emisses a DNA alexanders at MACE and M	RC6-UM0014-170400-023-G01 UM0014 Homo sentens cDNA	Homo septens X28 region near ALD tocus containing dual specificity phosphatase 9 (DUSP9), ribosomal protein L18a (RPL18a), Cat-Vicalmodulin-dependent protein kinase I (CAMKI), creatine transporter (CRTR), CDM protein (CDM) advandant contained to the contained of the contained o	Homo sepiens X28 region neer ALD locus containing dual specificity phosphatase 9 (DUSP9), ribosomal protein L18a (RPL18a), C22+/Calmodulin-dependent protein kinase I (CAMKI), creatine transporter (CRTR), CDM minden (CRTR),	Homo saplens polyglutamine-containing C14ORF4 gene
ZON FIODES	Top Hit Defebese Source	EST HIMAN	EST HIMAN	L	EST LIMANI	TA LINE	IN	Į.	EST HUMAN	L	¥	EST HIMAN	EST HUMAN	EST HIMAN	SWISSPROT	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	<u> </u>	T HIMAN	Т	Т			N
סוונלום ד	Top Hit Acession No.	BE268057 1		AL161491.2	R717941	AJ297357.1	AF147449.2	Γ	-	Γ	AJ131016.1		-	T		39.1		-	Γ		Γ	AV708305.1	U33472.1	2.	T				4
	Most Simitar (Top) Hit BLAST E Value	5.0E-03	5.0E-03	5.0E-03	5.0E-03	5.0E-03	5.0E-03	5.0E-03	5.0E-03	€.0E-03	5.0E-03	5.0E-03		4.0E-03			4.0E-03	-		4.0E-03	4.0E-03	4.0E-03	4.0E-03	-			4.0E-03	4.0E-03	
	Expression Signal	0.69	3.89	2.71	1.36	0.75	3.67	0.67	1.7	99.0	0.78	1.55	2.13	1.88	0.67	3.1	4.7	2.96	24.08	1.42	1.02	1.29	1.80	7.06	1.43	1.14	1.56	1.56	2.52
	ORF SEQ ID NO:	12905	13094		13126			13681			14445	14557	10304	10383	10495	10628	10927		11169	11188	11325		11775	12058		12314	12680	12581	12691
	Exon SEQ ID NO:	7885		8097	8109	Ш					9466	9568	6295	5374	5477	5629	5884	5918	6138	6155	6283	6548	6699	6953	7165	7192	7466	7466	7581
	Probe SEQ ID NO:	2865	3063	3081	3083	3204	3616	3673	3864	4179	4476	4580	232	319	440	869	868	900	1133	1151	1284	1551	1704	1968	2186	2215	2498	2498	2819

Page 56 of 209 Table 4

		_	Т	7	т	7	$\tau$	_	7	_		_	7	_	<b>-</b>	_	_	_	_	O		T	4	U		_	E	1 "		П	П	
Single Exon Probes Expressed in HBL100 Cells	Top Hit Descriptor		Indmo septens polygit/damine-containing C14ORF4 gene	Hamb septens chromosome 21 segment HS21C084	FM1-11 U340-151289-003-h08 HT0340 Homo saplens cDNA	PM1-H10340-151299-003-h08 HT0340 Homo sapiens cDNA	Application of Code Homo seplens cDNA clone IMAGE:2865279 3	Apontation NCI_CGAP_Cot8 Home sapiens aDNA clone IMAGE:2865279 3'	Home series Thints age 447	ab18a08.x5 Stratagene lung (#937210) Homo sapiens cDNA done IMAGE:841142.3' etniliar to contains Aku	Home eminer;	month september where CKZ catalytic subunit eigher gene, exch 1	norino seprensi protein kinase CRZ catalytic subunit alphe gene, exon 1 nc73c05.s1 NCI_CGAP_P72 Homo sapiens cDNA clone IMAGE:782984 similar to contains Als repetitive	omiral, (a. 11-1). This	A.c. cate (cv. nao) mrtvA for trosephosphale isomerase	Miss misseauus intesanai tretoi tactor gene, partial cols	Mus musculus intestina trefoi factor gene, partial cds Arehidoscia their	Analysis utiliana ipowi gene	00 LOJ 802FT NIH MGC 44 Homo saplens cDNA clone IMAGE:3608833 6*	Mus musculus alnea (VX/III) colloca (VXXIII)	Celegans same cone	AV762392 MDS Home septens cDNA close MDSBS-504 F	AV762392 MIDS Homo saplens cDNA clone MIDSBSG01 F	ah04f09.y5 Gessler Wilms tumor Home sapiens cDNA clone IMAGE-14 ggaso gr	S.cereale (cv. Halo) mRNA for triosephosphate Isomerase	Rattus novegicus gdirf gene	ht68g08.x1 NCI_CGAP_Lu24 Homo sepiens pDNA close MAAGE 3154281.31	xu8.P10.H3 conorm Homo septens cDNA 3'	ab18a08.x5 Stratagene lung (#637210) Homo saplens cDNA clone IMAGE:841142.5' similar to contains Alu	801482715F1 NIH MGC 68 Home emile - DNA -1. 1131 AF ASSESSED	Homo sapiens RAP1 GTPssa activation and the CAN EDIA	Homo saplens RAP1, GTP ase activating protein 1 (RAP1GA1) mRNA
Sedon Probes	Top Hit Deterbase Source	15/2	EN N	COT LAINAN	TO LOS	EST LINAN	EST LIMAN	SWISSPROT	LN		TN	12	5	PANOL 12	12	Į.	Į	EST LIMIAN	EST HIMAN	Z	Z	EST_HUMAN	ST_HUMAN	EST_HUMAN	LN	F	EST_HUMAN	П	EST HIMAN	Т		
	Top Hit Acession No.	A 1277388 4		BE154134 4	RE454494 4	AW188428 1	AW188428 4	013608	AJ011712.1	AI732754 1	AF011920.1	AF011920 4	AA468110 1	Z32521 1	U46858 1	U46858 1	Ī	Ţ	L	Γ	Y12500.1			1			1	41536141.1 E	3.0E-03 AI732764.1		4506414	4506414 NT
Most Similar	(Top) Hit BLASTE Value	4.0F-03	4.0F-03	4.0E-03	4 0F-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03	4.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03	3.0E-03 B	3.0E-03	3.0E-03
	Expression Signal	2.62	19.	1.06	1.06	0.92	0.92	1.02	1.73	1.08	2.62	5.77	272	7.09	0.92	0.92	0.83	4.48	3.09	2.08	7.49	7.05	7.05	1.6	1.04	4.26	0.68	5.15	2.19	8.19	0.92	0.92
	ORF SEQ ID NO:	12692		13191			13487			14451	10434	10928	11684		12323	12324		13038	13108	13361		13876	13877	13915		14254		14359	14663	14673	14922	14923
,	SEQ ID NO:	7581	7585	8170	8170	8460			8889	9470	5419	5886	6618	7208	7209	7209	7942	8028	8094	8343	8351	8871	8 8	C7/62	223	8 S	8328	8378	1298	9690	8945	8945
4	SEQ ID NO:	2619	2824	3154	3154	3452	3452	3815	3889	4480	88	898	1621	2231	2232	2232	2923	3008	3078	3333	3342	3870	2000	2500	/704	1/24	32	) (1)	4686	4705	4968	4968

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	CD11b=leukcoyte Integrin alpha chain (human, Genomic, 104 nt, segment 23 of 31)	RING CANAL PROTEIN (KELCH PROTEIN)	RING CANAL PROTEIN (KELCH PROTEIN)	yd15h03.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:108341 5	Human alpha-2-plasmin inhibitor gene, exons 6 and 7	nu86f01.s1 NCI_CGAP_Alv1 Homo saplens cDNA clone IMAGE:1217563	Homo saplens tumor-related protein DRC2 (DRC2) gene, complete cds	PLATELET-ENDOTHELIAL TETRASPAN ANTIGEN 3 (PETA-3) (GP27) (MEMBRANE GLYCOPROTEIN   SFA-1) (CD151 ANTIGEN)	Homo sapiens procollagen-lysine, 2-conglutarate 5-dioxygenase (lysine hydroxylase, Ehlers-Danks syndrome	type VI) (PLOD) mRNA	Homo sepiens procellagen-lystne, 2-cocglutarate 5-dloxygenase (fysine hydroxylase, Ehlere-Danios syndrome type VI) (PLOD) mRNA	COLLAGEN ALPHA 6(IV) CHAIN PRECURSOR	2x42a10.11 Scares_total_fetus_Nb2HF8_9w Homo explens cDNA clane IMAGE:789114 5	Mus musculus myelin expression factor-3-like protein gene, partial cds	Homo sapiens chromosome 21 segment HS21C102	UHH-BI1-adi-g-10-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2717010 3'	zx42a10.r1 Soares_total_fetus_Nb2HF8_9w Homo sepiens cDNA clone IMAGE:789114.5	602183960T1 NIH_MGC_42 Hamo sepiens cDNA clone IMAGE:4300070 3'	H.sepiens DMA, DMB, HLA-Z1, IPP2, LMP2, TAP1, LMP7, TAP2, DOB, DQB2 and RING8, 9, 13 and 14	garies  ENV DO VODOTEIN (CONTAINS: COAT DOOTEIN CORP. COAT DOOTEIN COAS)	Rathis novembring Scholandamine? recents need benfalleds	HILH-RWO-eing-0-11 at NCI CGAP Sub6 Homo septens cDNA close IMAGE:2730413.31	HA0507 Human fetal liver cDNA library Homo sapiens cDNA	Drosophila melanogaster shortsighted class 2 (shs) mRNA, complete ods	Drosophile melanogaster shortsighted class 2 (shs) mRNA, complete cds	yo45e02.s1 Soares #dult brain N2b4HB55Y Homo sapiens cDNA clone IMAGE:180890 3'	Homo saplens X-linked anhidratic ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat	subject	- 1	as70b08x1 Bersteed colon HPLRB7 Homo explens cDNA clone IMAGE:2334039 3' similar to TR:Q13825   Q13825 AU-BINDING PROTEIN/ENOYL-COA HYDRATASE.;	
Top Hit Database Source		SWISSPROT	SWISSPROT	EST_HUMAN	NT	EST_HUMAN	NT	SWISSPROT		T,	<b>-</b>	SWISSPROT	EST_HUMAN	NT	NT	EST HUMAN	EST HUMAN	EST HUMAN	_,	TOUGOUNG.	TA PER LA PERPETRA PER LA PERPETRA PER LA PERPETRA PERPETRA PER LA PERPETRA P	ECT LIMAN	EST HUMAN	Į.	Z	EST_HUMAN	 	¥	EST HUMAN	EST HUMAN	İ
rop Hit Acession No.	S52213.1			1		1.1		48500		4557836 NT	4657836 NT		38.1	F302691.1	L163302.2		2.0E-03 AA450138.1 E			8/344.1	103374 S	ļ	T					2	H96471.1	A1720263.1	
Most Similar (Top) Hit BLAST E Value	3.0E-03	2.0E-03 Q04852	2.0E-03 Q04652	2.0E-03	2.0E-03 M20783.1	2.0E-03 A	2.0E-03 A	2 0F-03 P		20E-03	2.0E-03	2.0E-03 P29400	2.0E-03/	2.0E-03 A	2.0E-03 /	2.0E-03/	2.0E-03	2.0E-03 B			2.0E-03	205.02	2.0E-03	2 0E-03	2.0E-03	2.0E-03		20E-03	1.0E-03	1.0E-03	
Expression Signal	98.0	0.69	0.69	10.87	1.92	1.98	8.74	4 39		1.7	1.7	4.37	1.16	1.19	1.03	3.15	5.57	0.86	,	19.6	2.35	3 .	0 92	222	222	6,		0.04	1.26	1.31	
ORF SEQ ID NO:	14961	10550	10551		11394	11398	11404	44544		11537	11538		11806	12036	12287		13360	13365			13980	1	14285	l	1				10490	10878	
Exan SEQ ID NO:	9885	5546	6546	7731	6343	6345	6354	6455		6482	6482	6553	6729	6935	7167	7472	8342	8347		8288	8993	300	0000	0405	9405	9561		9842	5474	6839	
Probe SEQ ID NO:	5014	544	511	111	1346	1348	1357	1458		1485	1485	1556	173	1949	2188	2504	3332	3338		3582	3997	800	4302	4445	4415	4573		4861	436	819	

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Table 4
Single Exon Probes Expressed in HBL100

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	Top Hit Descriptor	as70b08.x1 Barstead odon HPLRB7 Homo septens oDNA clone IMAGE:2334039 3' similar to TR:Q13825 Q13825 AU-BINDING PROTEIM/FNOYI -COA HYTRATASIF .	wk86a06.xt NCI CGAP Part Homo septems cDNA clone INACE: 2422258 3	wx83e10.x1 NCL CGAP_Mel15 Homo saciens cDNA clone IMAGE-2651242.3*	wd88a01.xf NG_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:2338440 3' similar to contains Alu receditive element:	HIGH MOLECULAR WEIGHT FORM OF MYOSIN (7HMWMI)	Homo sapiens SCL gene locus	Homo saplens mRNA for KIAA1291 protein, partial cds	CARBÓNIC ANHYDRASE VI PRECURSOR (CARBONATE DEHYDRATASE VI) (CA-VI) (SECRETED CARBONIC ANHYDRASE) (SALIVARY CARBONIC ANHYDRASE)	CARBONIC ANHYDRASE VI PRECURSOR (CARBONATE DEHYDRATASE VI) (CA-VI) (SECRETED CARBONIC ANHYDRASE) (SAI IVARY CARBONIC ANHYDRASE)	LINE-1 REVERSE TRANSCRIPTASE HOMOLOGO	Human MUC2 gene, promoter region	Human MUC2 gene, promoter region	Homo saplens SVMT gene for synaptic vesicle monoamine transporter worns 14 16	S.cerevisiae chromosome X reading frame ORF YJR149w	RC1-TN0128-160800-021-001 TN0128 Homo sapiens cDNA	TCBAP1D4909 Pediatric pre-B cell acute lymphoblastic leukamia Baylor-HGSC project≔TCBA Homo sapiens cDNA clone TCBAP4809	Ceenorhabditis elegans spliced leader RNA (SL3 alpha). (SL4), and (SL5) comes	ov45c04.x1 Soares_testis_NHT Homo saplens cDNA cione IMAGE:1640262.3'	ow45c04.x1 Sogres_tests_NHT Hamo septens cDNA clone IMAGE:1640262.31	PM0-HT0339-200400-010-D02 HT0339 Homo sapiens cDNA	APOLIPOPROTEIN A-IV PRECURSOR (APO-AIV)	ai31c12.s1 Soares_bests_NHT Homo sepiens cDNA clone 1376318 3' similar to SW.AATC_CHICK P00504 ASPARTATE AMINOTRANSFERASE_CYTOPLASMIC:	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG	Homo saplens prion protein (PrP) gene, complete cds	Home sapiens prion protein (PrP) gene, complete cds	Homo saplens chromosome 21 segment HS21C010	Homo sapiens chromosome X open reading frame 6 (CXORF6) mRNA	wj15a11.x1 NCI_CGAP_Kld12 Homo saplens cDNA clone IMAGE:24028763"
	Top Hit Database Source	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	SWISSPROT	L'N	LN	SWISSPROT	SWISSPROT	SWISSPROT	Z	Ę	FN	LN LN	EST_HUMAN	EST HUMAN	NT	EST HUMAN	EST HUMAN	EST_HUMAN	SWISSPROT	EST HUMAN	SWISSPROT	N F	Ę	N N	NT	EST_HUMAN
	Top Hit Acessian No.	Al720263.1			Al692616.1	P47808	AJ131016.1	AB033117.1	P18915	P18915		-	U68061.1	AB044400.1		BE939162.1	BE246536.1		_	AI073485.1		046409	AA815400.1		-	U29185.1	AL163210.2	185170	Al862525.1
\[ \]	Most Similar (Top) Hit BLAST E Value	1.0E-03			1.0E-03				1.0E-03	1.0E-03	1.0E-03		1.0E-03	1.0E-03	1.0E-03	1.0E-03	1.0E-03	-	1.0E-03	1.0E-03		1.0E-03	9.0E-04	-	-				6.0E-04
	Expression Signal	1.31	3.35	1.17	4.08	2.86	4.54	1.8	2.17	2.17	0.75	0.78	0.76	1.51	0.76	5.29	4.94	16.0	1.69	1.69	5.57	8.4	1.32	4.35	2.42	1.75	1.09	1.03	1.51
	ORF SEQ ID NO:	10877	11114	11135	11184	12069	12188	12949	13163	13154	13265		13489		13840	14288	14321	14505	14644	14645		14855	14843		14592	12430	12715		13860
	Exan SEQ ID NO:			6105	6152	6962	7074	7930	8134	8134	8243	8474	8474	8288	8833	9304	9340	9518	9862	9662	9663	9884	9875	9050	9604	7309	7602	8222	8855
	Probe SEQ ID NO:	819	1078	1098	1148	1977	2033	2911	3118	3118	3228	3466	3466	3581	3831	4312	4349	4528	4677	4677	4678	4905	4896	4056	4619	2335	2642	3207	3853

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	in the second se	Hamo Bapiens CARS diemarne receptar (Uninterno) gaire, withhere we	HYPOTHETICAL 29.3 KD PROTEIN (UR192)	QV0-CT0225-021099-030-030 CT0225 Homo septems cDNA	nk27e11.s1 NC_CGAP_Co11 Home sapiens cDNA clone IMAGE:1014764 3' similar to contains Alu	repeanve clearent,	Haemophilus influenzae Kd section 63 of 163 of the complete genome	as70b08.x1 Barstead colon HPLRB7 Hamo septens cDNA clane IMAGE-2334039 3' stritter to 1 K.Q.13623 Q13825 AU-BINDING PROTEINENOYL-COA HYDRATASE.;	as70b08.x1 Barstead colon HPLRB7 Homo saplens cDNA clone IMAGE:2334039 3' similar to TR:013825	Q13825 AU-BINDING PROTEINENOYL-COA HYDRATASE.;	RC3-CT0254-130100-023-f01 C10254 Hamb sapiens GUNA	Homo septens chromosome 21 segment H521 CU/8	DKFZp434D059_r1 434 (synonym: htes3) Homo sapiens cDNA cione DKrzp434D038 p	SERICIN-2 (SILK GUM PROTEIN 2)	Homo sapiens neuropilin 2 (NRP2) gene, complete cds, alternatively spliced	Homo sapiens chromosome 21 segment HS21C067	INTIDATO, ST NCI CGAP COT Hamo sapiens CDNA done IMAGE:951930 3' similar to gb:M21121 T-CELL	SPECIFIC RAINTES PROTEIN PRECURSOR (HUMAN);	Inh10a10.s1 NCI_CGAP_Co1 Homo septens cDNA clone IMAGE:861830 3' similar to go:MZ1121 1-CELL	SPECIFIC RANTES PROTEIN PRECURSOR (HUMAN);	Zn61c08.s1 Stratagene musicle 837.209 rigino sapiens culty didio minori	601345895F1 NIH MGC 8 Hamo sapiens curva cione invadeor/es tv o	ly/78510.s1 Soares_multiple_sderosis_znantwor normo explains cours cours invocations of contains. At repetitive element.	DKFZp761,J221_r1 761 (synonym: hemy2) Homo septens cDNA clone DKFZp761,J221 5	180 KD SECRETORY PHOSPHOLIPASE AZ RECEPTOR PRECURSOR (PLAZ-R)	Human short chain acyl CoA dehydrogenase gene, exons 1 and 2	azzedos. vi NCI CGAP Kid11 Homo saplens cDNA clone IMAGE:2028197 6	Hr23a02.x1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:2119082 3'	INTERNALIN B PRECURSOR	GLUTAMATE DEHYDROGENASE 2 PRECURSOR (GDH)	Home seniens Xa assudesubsomal region; segment 1/2	BCO HT0014 310509-078 HT0014 Homo sections CDNA	Т	1	٠
Top Hit Detabase Source		노	SWISSPROT	EST_HUMAN		EST HUMAN	NT	EST HUMAN		EST HUMAN	EST_HUMAN	Ę	EST_HUMAN	SWISSPROT	Į,	L		EST HUMAN		EST HUMAN	EST HUMAN	EST_HUMAN	COT CIMAN	EST HUMAN	SWISSPROT	ΙZ	EST HIMAN	EST HUMAN	SWISSPROT	TORGRENA		4444111	NAMOR I CE	EST HUMAN	
Top Hit Acession No.		U45983.1	5.0E-04 010341	AW851844.1		AA548831.1	4.0E-04 U32748.1	AI720263.1		4.0E-04 AI720283.1	4.0E-04 AW763358.1	4.0E-04 AL163278.2	AL046704.1	4.0E-04   O96615	4.0E-04 AF281074.1	4 0F-04 At 163267 2		4.0E-04 AA578331.1		4.0E-04 AA576331.1	4.0E-04 AA086324.1	BE560660.1	4 000 000 000 000 000 000 000 000 000 0	4.0E-04 1440013.1	9 0F-04 P49259	1 182001 1		3.0E-04 AI399674 1	9 0E-04 P25147	077070	-13	ין ד	~	4 BE153778.1	
Most Similar (Top) Hit BLAST E	8	6.0E-04 U4	5.0E-04	6.0E-04		5.0E-04	4.0E-04	4.0E-04 Ai		4.0E-04	4.0E-04	4.0E-04	4.0E-04	4.0E-04	4.0E-04	A OF DA	4.00.01										1							3.0E-04	
Expression Signel		3.01	10.72	4:		121	1.07	134		1.34	2.18	1.19	0.94	2.01	3.3	4 42	1.16	97.6		2.79	2.39	3.42		7.02				1.08				1.31	1.09	5.1	
ORF SEQ ID NO:		14041	10875			13357	10700	40896		10897	11491			12636		L		14160		14170	14382	14867		14963				11884			13865	6	*	8	
Exan SEQ ID NO:		9054	1	l	١.	8338	5691		1	5856	85. 45.25		L			l	8//9	9480		9189										1				1 9656	
Probe SEQ ID		4060	643	1472		3329	885	633	3	837	1437	2030	2075	2554	3084		3//6	4406		4196	4407	4915		5019	2	<u>8</u>	88	1803	101	3236	3857	3942	3976	4671	

Page 60 of 209 Table 4 Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Homo sapiens SCG10 like-protein, helicase-like protein NHL, M68, and ADP-ribosylation factor related protein 1 (ARFRP1) genes, complete cds	AU146707 HEMBB1 Homo sapiens cDNA clone HEMBB1001253 3	Human dystrophin gene	Human dystrophin gene	qh98e11.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1865052.3° similar to contains MER3.b2 MER3 repetitive element;	Homo saplens chromosome 21 segment HS21C003	Mus musculus 5' flanking region of PibS gene	21.39b05.s1 Sceres overy tumor NbHOT Homo sepiens cDNA clone IMAGE:740337 3' similar to contains Atu repetitive element.	Human germline T-cell receptor beta chain TCRBV17S1A1T, TCRBV2S1, TCRBV10S1P, TCRBV2BS1P, TCRBV16S1P, TCRBV16S1, TCRBV14S1, TCRBV16S1P, TCRBV3S1P, TCRBV3S1P, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S1, TCRBV3S1, TCRBV3S1, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S1, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S1, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S1, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S1, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S1, TCRBV3S1, TCRBV3S1, TCRBV3S2, TCRBV3S1, TCRBV3S1, TCRBV3S1, TCRBV3S2, TCRBV3S1	am58c09.xt Johnston frontal cortex Hamo sapiens cDNA clone IMAGE:1639760 3'	Homo sapiens tubulin, beta, 4 (TUBB4) mRNA	QV2-BT0636-070500-194-b07 BT0636 Homo sapiens cDNA	Human tyrosine kinase TXK (bX) gene, exons 9 and 10	ES 1380550 MAGE resequences, IMAGE from septens with	Phaseous Yugars mirate reductase (FYNYK) gens, whipped was	M01611.11 Scares pineal gland_North or name septents contraction invoces 22000 o	WUTSTILL SOZIES DIRECTORING INSTITUTION SQUARES COLOR INVOLVENCE OF THE COLOR	Gallus gallus processome zo Kus subulu mollowy illing of whiteon out	Ushio reno nagoramo gene, exons 1 to 9, partial cus	yz26c09.s1 Soares melanocyle zivorim nomo sapiens curva ciute invade zivorim commine in	RETROVIRUS-RELATED POL POLYPROTEIN (CONTAINS: REVERSE TRANSCRIPTASE;   IENDONLO: EASE]	III.H.RiG.ash.e.09-0-U.st NCI CGAP Sub1 Homo septems cDNA done IMAGE:2708825 3'	THE DISCONDING A NOT COAP Subt Homo saplens CDNA clone IMAGE:2708825 3'	Annual Annual dynamine D141 recents (4141) gene, complete cds	Angular digular department of the second sec
Top Hit Database Source	NT	EST HUMAN	NT	NT	EST_HUMAN	NT	IN	EST_HUMAN	¥	EST_HUMAN	Ę	EST_HUMAN	LN L	EST HUMAN	N	EST HUMAN	EST_HUMAN	EX.	LN	EST_HUMAN	POGGENA	TOTAL TOTAL	NAME I SEL	ESI DOMAIN	Z
Top Hit Acession No.	2.0E-04 AF217798.1 N		18524.1	36524.1	286021.1	L163203.2			U66061.1	11124529.1	5174736	3E082317.1	J34374.1	AW978441.1	U01029.1	H96265.1	H96265.1	U09226.1	AB037997.1	HB9646.1	044980	T	AW013847.1	AW013847.1	U62918.1
Most Similar (Top) Hit BLAST E Value	2.0E-04	2.0E-04	2.0E-04	2.0E-04 ME	2.0E-04 A	2.0E-04	2.0E-04	2.0E-04	2.0E-04	2.0E-04	2.0E-04	2.0E-04							2.0E-04	1.0E-04					1.0E-04
Expression Signal	1.62	6.11	8.36	6.36	3.83	207	0.97	1.24	3.16	1.18	1.1	2.67						1.39	1.87	1.09				"	3.3
ORF SEQ ID NO:	10260		10953	l					12583	L							14512		14838	10806				11134	
Exan SEQ ID NO:	6239	6511	5914	5914	6164	6170	L	<u> </u>		l	1_	1_		8818	9016			9636	9870	67.79					6309
Probe SEQ ID NO:	175	475	898	898	1180	1167	1796	22	2800	2918	3260	3351	3381	3816	4020	4534	4534	4651	4891	758		1058	1097	1097	1312

Page 61 of 209 Table 4 obes Expressed in HBL100 Cells

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Single Exon Probes Expressed in HBL100 Cells	Top Hit Descriptor	Kaposi's sarcoma-associated herpesvirus ORF 68 gene, partial cds; and ORF 69, kaposh, v-PLIP, v-cyclin, latent nuclear antigen, ORF K14, v-GPCR, putative phosphoribosyfformylglycinamidine synthase, and LAMP (LAMP) genes, complete ods	Kaposi's sarcoma-associated herpesvirus ORF 88 gene, partial cds; and ORF 89, kaposin, v-FLIP, v-cyclin, latent nuclear antigen, ORF K14, v-GPCR, putative phosphoribosyfromylglycinamidine synthase, and LAMP AMP cancer, complain cds.	Equus caballus DNA, chromosome 24q14, microsatellite TKY36	SPLICEOSOME ASSOCIATED PROTEIN 62 (SAP 62) (SPLICING FACTOR 3A SUBUNIT 2) (SF3A66)	ij01f11.xt NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2140269 3' straliar to contains Atu repetitive dement;	Mouse alpha 1 type-IV collagen mRNA	AV647727 GLC Homo sapiens cDNA done GLCBBD043'	Homo sapiens KIAA0237 gene product (KIAA0237), mRNA	Homo sepiens KIAA0237 gene product (KIAA0237), mRNA	ah45c11.s1 Soares_testis_NHT Homo sapiens cDNA clone 1282468 3'	Pisum sativum mRNA for beta-1,3 glucanase (gns2 gene)	Pisum sativum mRNA for beta-1,3 glucenase (gns2 gene)	wy78e04,x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo septens cDNA clone IMAGE:2554638 3	RC3-CT0208-220999-011-E04 CT0208 Homo sepiens cDNA	RC3-C10208-220999-011-E04 CT0208 Homo saplens cDINA	HUM072014F Human foves cDNA Homo saplens cDNA clone EST HrD072014	HUM072014F Human fovea cDNA Homo sapiens cDNA clone ES1 MrD072014	PROBABLE GLYCEROL-3-PHOSPHATE ACYLTRANSFERASE, MITOCHONDRIAL PRECURSOR (GPAT)	Homo sapiens chromosome 21 segment HS21C078	Dictyostellum discoldeum gene for TRFA, complete cds	lg73c08.x1 Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGE:2114416.3	Homo saplens jun dimertzation profein gene, partial cds; cfos gene, complete cds; and unknown gene	Homo sapiens chromosome 21 segment HS21C001	Rat cytomegalovirus Maastricht, complete genome	Homo saplens chromosome X open reading frame 6 (CXORF6) mRNA		
xon Probes	Top Hit Database Source	ŢN		 	SWISSPROT	EST HUMAN	N	EST_HUMAN	NT	NT	EST HUMAN	NT	NT	<b>EST_HUMAN</b>	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	SWISSPROT	IN	N.	EST_HUMAN	<u> </u>	۲N	NT	TN	NT	
Single E	Top Hit Acession No.	48805.1	4000E 4	Τ	1.0E-04 Q62203	1.0E-04 AI440282.1		1.0E-04 AV647727.1	7662015 NT	7862015 NT	AA718833.1	8.0E-05 AJ251646.1	8.0E-05 AJ251646.1	AW044605.1	AW847445.1	AW847445.1	7.0E-05 L49075.1	7.0E-05 L49075.1	022949	7.0E-05 AL 163278.2	AB009080.1	7.0E-05 Al432413.1	7.0E-05 AF111167.2	7.0E-05 AL163201.2	9845300 NT	4885170 NT	4885170 NT	
	Most Similar (Top) Hit BLAST E Vatue	1.0E-04 AF1	, d	1.0E-04 AB	1.08-04	1.0E-04	1.0E-04	1.0E-04	1.0E-04	1.0E-04	9.0E-05 AA7		8.0E-05	8.0E-05 AW					Ì	ı	L			L				
	Expression Signal	3.97		1.7	0.96	0.72	1.86	1.09	1.64	1.64	1.84	1.19	12.78	0.67	11.8	11.8	1.24	1.24					0.72					
	ORF SEQ ID NO:	11646		11908		13863								14323		10405	10595	10596		L	\  -			14225	L			]
	SEQ ID NO:	6585		6814		8657	L						L	L		<u> </u>			L	ı			8041	L	L	Ł	1_	
	Probe SEQ ID NO:	1588		1556	3211	3651	3948	3867	4925	4925	689	811	853	4353	345	345	562	582	1038	2846	3085	3617	3043	4247	4760	1078	1976	

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	wb54h08.x1 NCI_CQAP_GC8 Homo sapiens cDNA clone INAGE:2308631 3' similer to gb:J03250 DNA TOPOISOMERASE I (HUMAN);	Homo sapiens monocyte/neutrophil elastase inhibitor gene, complete cds	QV4-ST0234-241199-040-h11 ST0234 Hamo eaplens cDNA	Homo sapiens 22kDa percetsomal membrane protein-like (LOC55885), mKNA	Homo saplens MEP1A gene, promoter region and exon 1	Homo sapiens partial SLC22A3 gene for extremeuronal monoamime transporter (EM 1), extra 1	Human renin (REN) gene, 5 flanking region	RETINAL-BINDING PROTEIN (RALBP)	RETINAL-BINDING PROTEIN (RALBP)	Cryptosporidium parvum Isolate Zaire 15 KDe grycoprotein gp10 gene, peruti cus	qh64c10x1 Soares fetal liver splear 1NFLS S1 Homo sapiens cDNA cione IMAGE:1849456 3 similar to	Contains Alu repetitive etement contains agement NEX repouted exercise.	W24g03X1 Soares WFL 1 GGC 31 name suppers convious involved and a second size	601461463F1 NIH MGC 00 HOMO septems curve cross involves constructions	601461463F1 NIH MGC 06 Homo sapiens cumA cione image. Jood 142 9	SKELEMIN	PM1-H10527-120200-001-010 H10521 H0mio septens conv.	PM1-H10521-120200-001-e10 H10521 Homo segions control	EST79996 Placenta I Homo saplens CLINA similar to similar to pos-associated protein	EST78896 Piacenta I Homo sapiens CUNA similar to similar to progressionation progress	Homo sapiens NOD1 protein (NOD1) gene, excits 1, 2, 4tha 3	AU125/21 N I ZKM4 Homo sapiens GUNA Gone N I ZAWHOUZOVO O	qh98e11.x1 Soares_NFL_T_GBC_S1 Homo sepiens CDNA ciche IMAGE:1855002.5 entue to contents. MER3.b2 MER3 repetitive element;	Human adenosine deaminase (ADA) gene, complete cds	zq46a12.r1 Stratagene hNT, neuron (#937233) Homo sapiens cONA clone IMAGE:532734 6' similar to	contains Alu repetitive element contains element L1 repetitive element;	RC3-BT0319-120200-014-h08 BT0319 Home sapens curv.	Homo sapiens p47-pnox (NCr1) gene, complete cos	H.sapiens DNA for endogenous retroviral like diement	S.cerevisiae 12.8 Kbp fragment of the left arm of chromosome XV	Homo saplens chromosome 21 segment HS21C082	Drosophila melanogaster strain Lamto 120 Suppressor of Harress (Su(H)) gone, purusi ous	
	Top Hit Dertabase Source	EST HUMAN	ri .	EST_HUMAN	ri		<u></u>		SWISSPROT	SWISSPROT	Į.		EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	SWISSPROT	EST HUMAN	EST HUMAN	ST HUMAN	EST HUMAN	Z	EST HUMAN	EST HUMAN	N.	j	EST HUMAN	EST_HUMAN	N	Ł	TN.	Į,	NT	   
}	8	<u> </u>	둗	ES	8923891 NT	Ā	N	NT	S	S	Z	-	삤	쁴	ŭ	ŭ	5	ŭΙ	Ü	Ü	Ψ)	Z	9	Ш	F	<del> </del>	3	E	_	=	1	۴	٢	1
Digino	Top Hit Acession No.	6.0E-05 A1655241.1	6.0E-05 AF053830.1	5.0E-05 AW392088.1	89238	AJ251058.1	AJ251884.1	U12821.1	4.0E-05 P49193	P49193	4.0E-05 AF164488.1		3.0E-05 AI248061.1	3.0E-05 AW273851.1	3.0E-05 BF037898.1	BF037898.1	3.0E-05 Q62234	3.0E-05 BE169211.1	3.0E-05 BE169211.1	3.0E-05 AA368679.1	3.0E-05 AA368679.1	3.0E-05 AF149773.1	AU125721.1	2 0E-05 A1286021.1	M13792.1		-		AF184614.1	X89211.1				
	Most Similar (Top) Hit BLAST E Value	6.0E-05/	6.0E-05/	5.0E-05	5.0E-05	5.0E-05 AJ	5.0E-05 A	4.0E-05	4.0E-05	4.0E-05	4.0E-05		3.0E-05	3.0E-05	3.0E-05	3.0E-05	3.0E-05	3.0E-05	3.0E-05	3.0E-05	3.0E-05	3.0E-05	3.0E-05	2.05-05	2 OE-05		2.0E-05		2.0E-05		L			
	Expression Signal	1.65	2.86	65.3	2.39	0.67	404	3.87	0.92	0.92	66.0		0.69	1.01	1.64	1.64	0.92	9.42	9.42	0.84	0.94	0.76	0.93				6.26							
	ORF SEQ ID NO:	12507	10704	11424		12828	13878		14325	14326			10706	11082	11148	11149	12720	14237	14238		14317		14663					13095				12804		
	Exan SEQ ID NO:	1,847	5698	6376	6816	7811	8873	6289	8348	9346	9710		2697	8052	6120	6120	7608	9251	9251	9333	9333	9476	9680		200		7601	808	8284	9000	N S	2423	١	١
	Probe SEQ ID S NO:	28.3	2742	1379	1828	2780	3872	2735	4355	4355	4725		672	1042	1114	1114	2848	4257	4257	4342	4342	4486	4605		707	onez	2841	3084	3272		328	3414	700	2002

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Table 4
Single Exon Probes Expressed in HBL100 Cells

				_	_	_	_		_			T A	_	-	_	_	•	4	<del>"                                     </del>			7	Т	Т	٦	7	-	<b></b>
	Top Hit Descriptor	Homo sepiens calcium chernel alpha1E subunit (CACNA1E) gene, exxis / -4s, and pana cas, euaneavoy spiloed	MOSAIC PROTEIN L'AN	Homo sapiens chromosomia 41 segment in 24 toward	Zwogget, I Soales was 1, 24 4 Home seriens DNA clone IMAGE: 2856548 3'	XMS911.X I NOT COAP HSC3 Home septems cDNA clone IMAGE:2246386 3'		9g11b08.x1 Scares_placenta_8b9weeks_2NbHP8b9W Home sapiens cDNA clone IMAGE:1759191 3:	Charles and Monday And CT0283 Home saciens CDNA	Accepted a Straighton of (#837210) Homo sapiens aDNA clone IMAGE:854251 3' straight to contains	MEROIT MERO repetitive element; MEROIT MERO TATALORIS Element;	Homo sapiens KIAAU330 gene product (n.j.A.v.35), in a.v.,  Homo sapiens KIAAU330 gene product (n.j.A.v.35), in a.v.,	qw16g09x1 NCi_CGAr_Uta nonio sapitatis calex contaminates	DAY2-BT0379-010300-105-411 BT0379 Horo saplens cDNA	OVABIAN ABIINDANT MESSAGE PROTEIN (OAM PROTEIN)	OVA 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	OVABIAN ARIUNDANT MESSAGE PROTEIN (OAM PROTEIN)	Annaly of Spares feta liver spleen 1NFLS S1 Homo sepiens cDNA clone IMAGE:1655738 3' similar to	contains MERS. 2 MERS repetitive element; contains MERS. 2 MERS prepetitive element; contains MERS. 2 MERS prepetitive element;	repetitive element contains L1 repetitive element; repetitive element contains L1 repetitive element contains cont	repetitive element contains defined by the second of the s	masa-no vi NCI CGAP HSC2 Homo sapiens cDNA clone IMAGE:2056168 3'	00020000000000000000000000000000000000	11. H. Ringast 505-0-U. S1 NCI CGAP Sub1 Homo sapiens cDNA clone IMAGE:271042631	Gallus nallus Dach2 protein (Dach2) mRNA, complete cds	III 3_CT02/4-150200-074-B03 CT02/4 Homo saplens cDNA	WIGHOUT NOT COAP BIR25 Home explens CDNA clone IMAGE: 2432562 3' similar to contains element	MER22 repetitive element :
	Top Hit Database Source	NT	SWISSPROT	Į,	EST HUMAN	EST HUMAN	EST HOMAN	EST HUMAN	TN	EST HUMAN	EST HUMAN	FN	_	EST HUMAN	ESI HOMAN	SWISSPROI	ES! HUMAN	SWISSPRO	EST_HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	EST TOWAR	TOT TOTAL	ES - TOWN	NI TOT	LS L	EST HUMAN
Y alfulc	rop Hit Acessian No.	F223391.1	П				A 583811.1	-		AW362539.1	AA669729.1	7682177 NT		Al368252.1			BE069189.1	Q01456	A1040099.1	R16267.1					3 AW015401.1	S AF198349.1	3 AW848295.1	4.0E-06 A1886839.1
	Most Similar (Top) Hit BLAST E Value	1.05-05/	1.0E-05	1.0E-05/	1.0E-05	1.0E-05	9.0E-06		9.0E-08	8.0E-08	7.0E-08	7.0E-08		7.0E-06	6.0E-06	6.0E-06	6.0E-06	8.0E-06	6.0E-06	4.0E-08						4.0E-06		
	Expression Signal .	4	10.08	1.04	1.76	1.9	2.89	4.56	2.64	1.52	1.3	242		6.96	1.18	1.03	0.83	1.96	2.14	5.78	86.9						1.07	1.95
	ORF SEQ ID NO:		13867	14033	14132	14671	12681	13048		12543		1468			12883	12912	13625	12912	14587	10670		L	11360		1 12304	13021	13807	14635
	SEQ ID NO:		8882	8	9150	8898	7563	8039	8535	11111	F079	8410		7822	7867	7890	8616	7890	9801	5887	<u> </u>		١.	L	L	8008		9648
	Probe SEQ ID SOO:	1	3880	4049	4155	4703	2601	3022	3529	2458	790			2802	2847	2871	3609	4609	4616	639	838	1314	1314	1445	2203	2990	3789	4683

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51 ORGANIC CATIONICARNITINE TRANSPORTER 2 (SOLUTE CARRIER FAMILY 22, MEMBER 5) (HIGH-AFFINITY SODIUM-DEPENDENT CARNITINE COTRANSPORTER) va04s03.x1 NCI\_CGAP\_Kid11 Homo saplens cDNA clone IMAGE:2297068 3' similar to contains MER30.b1 Human ABL gene, excn. 1b and Intron. 1b, and putative M8504 Met protein (M8504 Met) gene, complete cds W22a05 x1 NCI\_CGAP\_Ut1 Homo septens cDNA clone IMAGE:2426616 3' similær to TR:060734 060734 LINE-1 LIKE PROTEIN ;contains L1.t2 L1 repetitive element; hq64d12x1 NCI\_CGAP\_HN13 Homo septens cDNA clone IMAGE:3124161 3' hq64d12x1 NCI\_CGAP\_HN13 Homo septens cDNA clone IMAGE:3124161 3' ak48g11.s1 Sogres\_testis\_NHT Homo explens cDNA clone IMAGE:1409252 3' similar to contains LTR1.13 234b08.s1 Soares\_fetal\_liver\_sploon\_1NFLS\_S1 Homo capiens cDNA clone IMAGE:432863 3' similar to 234b08.s1 Scares\_fetal\_liver\_spleen\_1NFLS\_S1 Hamo sepiens aDNA clane IMAGE:432883 3' similar to POL POLYPROTEIN (CONTAINS: PROTEASE; REVERSE TRANSCRIPTASE; ENDONUCLEASE) Homo sapiens gene for alpha-1-microglobulin-bikunin, exons 1-5 (encoding alpha-1-microglobulin, Nzp02e05.rf Stratagene ovarian cancer (#337219) Homo septens cDNA chore IMAGE:895232 6 Human glyceraldshyde-3-phosphate dehydrogenase (GAPDH) gene, complete ods HOMEOBOX PROTEIN GOOSECOID Human glycereldehyde-3-phosphate dehydrogenase (GAPDH) gene, complete ods KNOB-ASSOCIATED HISTIDINE-RICH PROTEIN PRECURSOR (KAHRP) Vus musculus DBMMSE protein (DBMm5e) mRNA, complete cds Top 14t Descriptor Mus musculus gene for odcrant receptor A16, complete cds QV0-ST0247-090200-105-c05 ST0247 Homo seplens cDNA AV657555 GLC Homo sapiens cDNA clone GLCFDB05 3' Homo saplens p47-phox (NCF1) gene, complete cds Homo septens p47-phox (NCF1) gene, complete cds HISTIDINE-RICH GLYCOPROTEIN PRECURSOR Homo sepiens chromosome 21 segment HS21C078 Homo sapiens PP1200 mRNA, complete cds contains L1.tf L1 repetitive element; contains L1.11 L1 repetitive element Single Exon Probes Expressed in HBL100 Cells **VER30** repetitive element; LTR1 repetitive element; terminus.) SWISSPROT SWISSPROT EST HUMAN EST HUMAN EST\_HUMAN EST HUMAN SWISSPROT EST\_HUMAN EST\_HUMAN SWISSPROT EST HUMAN SWISSPROT SWISSPROT EST\_HUMAN EST HUMAN EST\_HUMAN Top Hit Database Source Ę 눋 Ę 눋 Top Hit Acession BE047094.1 BE047094.1 1.0E-08 AF184614.1 4.0E-06 AW817268.1 2.0E-06 AA173518.1 2.0E-06 AB030896.1 1.0E-06 O76082 1.0E-06 AF084364.1 1.0E-06 AF184614.1 3.0E-06 AA700562.1 3.0E-06 AA700562.1 3.0E-06 AF202635.1 3.0E-08 AA868218.1 2.0E-06 AV657555. 호 3.0E-06 AIB57779.1 2.0E-06 AI672138. 1.0E-06 U07581.1 3.0E-06 X54816.1 2.0E-06 P04929 2.0E-06 P06719 **J04038.**1 1.0E-06 P09125 2.0E-06 P54366 2.0E-06 P21414 1.0E-06 3.0E-06 3.0E-06 3.0E-08 単(金) BLASTE 13.66 0.95 238 1.48 4.19 1.11 1.24 1.63 0.82 3,32 1.07 1.73 1.73 4.34 32 8 1.29 1.19 60: \$ Spresslon Signer 13475 10684 11481 11544 14221 10100 12034 12035 14406 13695 12185 12886 13701 14739 12577 12194 14987 ORF SEQ Ö N O 8238 7462 5114 9750 5266 8449 8681 6489 7080 9750 8698 8699 9420 6533 280 8693 587 6934 7080 SEQ ID 10018 7871 8207 옂 2000年 4244 2315 2397 3441 3676 3689 1491 1988 4766 4766 1535 1948 Probe SEQ ID 2080 2205 2851 3191 3695 3695 4430 5047 ÿ

Page 65 of 209 Table 4 Probes Expressed in HBL100 Cells

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Single Exon Probes Expressed in MBL100 Cells	Top Hit Descriptor	Homo saplens chromosome 21 segment HS21C085	Homo sapiens chromosome 21 segment HS21C085	Homo sapiens glypican 3 (GPC3) gene, partial cds and tlanking report regions	Homo sapiens glypican 3 (GPC3) gene, partial cds and flanking repeat regions	qi82g07.x1 Soares_NhHMPu_S1 Homo saplens cDNA clone IMACE:18/38/6 3	qi82g07.x1 Soares_NhHMPu_S1 Homo sapiens cDNA cione IMAGE:18788763	CM3-CT0277-221099-024-e11 CT0277 Homo sapiens cUNA	Homo sapiens HLA class ill region containing banascin X (banascin-X) gene, partial cds, cyrocitroma F4bu Z1- hydroxylase (CYP21B), complement component C4 (C4B) G11, helicase (SKI2W), RD, complement factor B	(Bf), and complement component C2 (C2) genes,>	HYPOTHETICAL 24.1 KD PROTEIN IN LEF4-P33 INTERGENIC REGION	wh64f10.x1 NCI_CGAP_Kid11 Hamo sapiens cDNA clane IMAGE:2385547 3	EST93615 Supt cells Homo sapiens cDNA 5' end	Homo sapiens NOD1 protein (NOD1) gene, exons 4 through 14 and complete ods	ws84h05.x1 NCI_CGAP_Co3 Homo sepiens cDNA clone IMAGE:2504697 3'	Human microfibril-associated glycoprotein (MFAP2) gene, putative promoter region and alternatively spliced	untranslated exons	Homo saplens Xq pseudoautosomal region; segment 1/2	Human polymorphic microsatellite DNA	Human IgK subgroup I germline gene, excris 1 and 2, V-region 018 aliele	Human polymorphic microsatellite DNA	M.Ro-BN0115-020300-001-f11 BN0115 Hamo sapiens cDNA	MR0-BN0115-020300-001-f11 BN0115 Hamo sapiens cDNA	yd50f12.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA come INAGE:1110950 0	HYPOTHETICAL 63.8 KD PROTEIN IN GUTT-RIM1 IN LENGENIC REGION PRECURSOR	AV650201 GLC Homo saplens cDNA done GLCCCD013	yc14h09.s1 Stratagene Iung (#337210) Homo sapiens cDNA clone IMAGE:50700 3 Similar to sumer to gab: M62982 ARACHIDONATE 12-LIPOXYGENASE (HUMAN)	yc14h09.s1 Stratagene king (#837210) Homo sapiens cDNA clone IMAGE:80705 3' similar to similar to conceptation.	Homo saplens TRF2-interacting telemente RAP1 protein (RAP1) mRNA, complete cds	Homo sapiens DiGeorge syndrome critical region, telomeric end	Homo saplens DiGeorge syndrome critical region, telomeric end	Firm rubrices beta-cytoplasmic/vascular) actin gene, complete cds	
n Probes L	Top Hit Database Source			_		EST HUMAN	EST_HUMAN	EST HUMAN		_	SWISSPROT	EST_HUMAN	EST_HUMAN		EST HUMAN			<u>.</u>	LI .	L.	ĭ	EST_HUMAN	EST_HUMAN	EST_HUMAN	SWISSPROT	EST_HUMAN	EST HUMAN	T H IMAN	L	TN	FN	F.V	_
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Single	Top Hit Acession No.	NL163285.2	1.0E-06 AL163285.2	9.0E-07 AF003529.1	8.0E-07 AF003529.1	41288596.1	AI288596.1	4W855558.1		6.0E-07 AF019413.1	P41479	5.0E-07 AI831893.1	AA380630.1	AF149774.1	4.0E-07 AW009602.1	- ,	U19719.1	3.0E-07 AJ271735.1	3.0E-07 M99149.1	3.0E-07 M64857.1	3.0E-07 M99149.1	3.0E-07 BE005077.1	3.0E-07 BE005077.1	184704.1	P38739	3.0E-07 AV650201.1	T67850 1	TR7950 4		2 0E 07 1 77560 4	177560 1	Z.UE-07 L.7 209.1	U30048.1
	Most Similar (Top) Hit BLAST E Value	1.0E-08	1.0E-08	9.0E-07	8.0E-07	8.0E-07	8.0E-07	6.0E-07		6.0E-07	6.0E-07 P4	5.0E-07	5.0E-07	6.0E-07	4.0E-07		3.0E-07 U1	3.0E-07	3.0E-07	3.0E-07	3.0E-07	3.0E-07	3.0E-07	3.0E-07	3.0E-07	3.0E-07	TIZU-BU E	7030 6	2000	1			_
	Expression Signal	124	1.24	1.02	1.02	5.07	5.07	2.73		2.45	1.89	6.0	1.83	1.28	1.86		5.45	2.79	1.32	1.79	0.94	26.28				8.15	4				23.04		129.68
	ORF SEQ ID NO:	14887	14888	10421	10422	14598	14599	11945		12513				14480	13892		10492	L	L		12320			L		14566	77077	_					10255
	SEQ ID	9910	9910	5409	5409	9610	9610	6857		7392	8867	5378	6050	9501	88833		6476	5810	6353	6583								1	1			6219	
	Probe SEQ ID NO:	4833	4933	359	320	4625	4625	1868		2421	3865	324	96	4511	3893		438	678	1356	1586	2224	288	2399	2964	3086	4587	1	4080	4 689 5	87	153	153	19

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Table 4
Single Exon Probes Expressed in HBL100 Cells

									_			_					J	21		Į	←	u	5	_	Д	Ľ	<b> -</b> -4	4	P		F	J
Single Exon Probes Expressed in ABL100 Cells	Top Hit Descriptor	Homo expiens homeobox protein CDX4 (CDX4) gene, complete cds and flanking repeat regions	Homo septens homeobox protein CDX4 (CDX4) gene, complete cds and transing repeat regions	208b07.s1 Stratagene NT2 neuronal precursor 937230 Homo sapiens cDNA clone IMAGE:650869 3" similar to gbt.31660 GLYCOPHORIN A PRECURSOR (HUMAN);contains Atu repetitive element;	yor 5g04.st Stratagene lung (#837210) Homo sapiens cDNA ctone IMAGE:80790 3' similar to contains L1 repetitive element;	16 AUTOANTIGEN	HYPOTHETICAL 72.6 KD PROTEIN C2F7.10 IN CHROMOSOME I	Homo sapiens caveolin 1 (CAV1) gene, exon 3 and partial cds	Homo sapiens chromosome 21 segment HS21C082	GLYCOPROTEIN GPV	Homo sapiens chromosome 21 segment HS210082	AV718682 GLC Homo saplens cDNA clone GLCFNFU4 o	AV718662 GLC Homo sapiens CDNA clone GLCFNFU4 D	2856g02.rf Soares retina N2D4HK Horino septems curve during invade. 3002.rf Soares retina n2D4 curve INAACE-320273 34	Wd16b05x1 Sogres NrL_I_cb0_31 numb septem controller co	601690133F1 NIH_MGC_/ Hamo septemb culva clane invacionaryo o	601580133F1 NIH MGC / Hans sapers corn dura mandered of	ANKYRIN 1 (ERYTHKOCT IE ANNTRIN)	Ket miking for indeemal protein LD i	DINEIN REAVI CHAIN (DINC)	United and the second of the second HS210048	Turno septembrane 21 septembrit HS21C048	THOSE LITTORES ADVISOR OF A TOTAL SERVICES CONA	MINCHI DISCOULD AND AND AND AND AND AND AND AND AND AN	Inditio Sapieris N. Provinci (1977-1974)	Inomo saprens cinomicsonia x 1 cognitati HS2/C4/C4/C3	Home septems chicumodina at septems and a clare IMAGE:843183 similar to contains Alu repetitive	inusbussi Nojecov jing i ising oprasi element	DORSAL-VENTRAL PATTERNING TOLLOID PROTEIN PRECURSOR	DORSAL-VENTRAL PATTERNING TOLLDID PROTEIN PRECONSOR	DKFZp434J0426_r1 434 (synonym: https3) Homo sapiens curva cione un representes o	
Hobes L	Top Hit Detaberse Source			EST HUMAN	EST HUMAN	SWISSPROT	SWISSPROT			SWISSPROT		EST_HUMAN	EST_HUMAN	EST HUMAN	T HUMAN	EST HUMAN	EST HUMAN	SWISSPROT		SWISSPROT	SWISSPROT		1400	ESI HUMAN		<u> </u>	Ę	EST HUMAN	SWISSPROT	WISSPROT	EST_HUMAN	
<u> </u>		토	Z	ES	, E		S	뉟	N	NS.	Ŋ	ES	S	Ш П	EST	낕	낕	5	틸	গ্ৰ	S)	<u>z  </u>	zΙ		<u>ک</u> و	킼	<del>-</del>	ш	S	8	Щ	
Single	Top Hit Acession No.	\F003530.1	2.0E-07 AF003530.1	AA223280.1	2 0E 07 TR3042 4	026768	209701	2.0E-07 AF125348.1	1.0E-07 AL163282.2	P09258	1.0E-07 AL163282.2	1.0E-07 AV718662.1	1.0E-07 AV718662.1	1.0E-07 AA019181.1	8.0E-08 AI911352.1	8.0E-08 BE795469.1	BE795469.1	7.0E-08 Q02357	7.0E-08 X04809.1	7.0E-08 P15305	7.0E-08 P15305	6.0E-08 AL163248.2	6.0E-08 AL163248.2	BE14439	7662473 NT	6.0E-08 AL163248.2	5.0E-08 AL163303.2	5 0F-08 AA493851.1	4.0E-08 P26723	P25723	AL079581.1	
	Most Similar (Top) Hit BLAST E Vatue	2.0E-07	2.0E-07	2.0E-07.AA	2.05.07	2.0E-07 103042	2.0E-07 Q09701	2.0E-07	1.0E-07	1.0E-07 P09258	1.0E-07	1.0E-07	1.0E-07	1.0E-07	8.0E-08	8.0E-08	8.0E-08	7.0E-08	7.0E-08	7.0E-08			6.0E-08									
	Expression Signal	1.48	1.46		8	80.1	2.13	20.05	1.43	2.14	3.91	2.86	2.86	1.22	2.23	8.0	1.7	3.2	42.98				2.78			0.92	2.15	89 +				
	ORF SEQ ID NO:	10785	10786	000	2000	10981	11625			11543		14150						10167	11392	13517		10858	10859	12397	13019	14106	_		44703	1		
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	Probe SEQ ID NO:	730	200	8	RZA	088	1140	3604	1085	2764	3659	4169	4169	4973	604	1033	3468	79	1344	3495	3495	8	807	2302	2988	4126	ន		27/4	1/2	77.77	Ş

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	x87f06x1 NCI_CGAP_Lu26 Hamo saplens cDNA clane IMAGE:27871393'	zw48f07.r1 Soares_total_fetus_Nb2HF8_9w Homo espiens cDNA clone IMAGE:773317 6' similar to contains Alu repetitive element:contains element MER15 repetitive element ;	Gallus gallus Dach2 protein (Dach2) mRNA, complete cds	MR0-OT0080-240200-001-g08 OT0080 Homo saplens cDNA	MR0-OT0080-240200-001-g08 OT0080 Homo septens cDNA	601155321F1 NIH_MGC_21 Homo saplens cDNA done IMAGE:3138883 6*	Homo saplens chromosome 21 segment HS21C047	601570463F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3845189 5'	xp43f11.x1 NCI_CGAP_HN11 Hamo sepiens cDNA clane IMAGE:2743149 3'	rw64h01.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1251408 3' samilar to contains L1.t3 L1 resettive element;	Sheep His-IRNA-GUG	WNT-14 PROTEIN PRECURSOR	WNT-14 PROTEIN PRECURSOR	RC3-ST0197-161099-012-b03 ST0197 Homo septens cDNA	aa28c07.r/ NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:814380 5' similar to contains L1.t2 L1	repeative element;	repetitive element;	Homo sapiens caveolin 1 (CAV1) gene, exon 3 and partial cds	PM2-HT0130-150999-001-f12 HT0130 Homo eaplens cDNA	TCBAP1D5232 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo	saplens cDNA clone TCBAP5232	TCBAP105232 Pediatric pre-B cell acute lymphoblastic leukemia baylor-higsic project=1 obst meno septens cDNs clone TCBAP5232	Homo sapiens chromosome 21 segment HS21C079	Homo saplens chromosome 21 segment HS210079	Home sentens DNA for 3-ketosov/-CoA thiolese bets-subunit of mitochondrial trifunctional protein, exon 2, 3		Homo sapiens jun dimertzation protein gene, partial cds; cfos gene, complete cds; and unknown gene	PM1-HT0527-160200-001-h05 HT0527 Hamo sapiens cUNA	RC2-HT0252-120200-014-h10 HT0252 Homo sapiens culva	
Top Hit Deterbase Source	EST_HUMAN	L HUMAN	¥	EST HUMAN	EST_HUMAN	EST_HUMAN		EST_HUMAN	EST HUMAN	EST HUMAN		SWISSPROT	SWISSPROT	EST HUMAN		EST HUMAN	EST HUMAN		EST HUMAN		EST HUMAN	EST HUMAN						EST HUMAN	EST_HUMAN	
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Top Hit Acession No.	2.0E-08 AW302998.1	2 DE-08 AA425598.1	2.0E-08 AF198349.1	20E-08 AW886438.1	2.0E-08 AW886438.1	2.0E-08 BE280477.1	2.0E-08 AL163247.2	BE734871.1	2.0E-08 AW270271.1	2 NE-OR 44731948 1	2 0F-08 K00216.1	042280	042280	AW813620.1		AA459040.1	AW572881.1	AF125348.1			BE246844.1	BE246844.1		_		D86842.1	AF111167.2	BE169421.1		
Most Similar (Top) Hit BLAST E Vatue	2.0E-08	2.05-08	20E-08	20E-08	2.0E-08	2.0E-08	2.0E-08			1		İ				2.0E-08	2.0F-08				1.0E-08			L		7.0E-09	6.0E-09	L		
Expression Signal	8.87	7.80	2.63	9.13	9.13	26.56	1.74	1.75	3,33	,	18.	A 45				1.32	3 44				1.06					1.62	1.19			
ORF SEQ ID NO:			10637				11369	L			2	424.72						77077			13165	l	١				14287			
Exen SEQ ID NO:	5270	£300	5529	5880	2680	5991	l			_	7444	2405 0464		ł		9274	7020	1			3119 8135	1	1	4120 8114 4120		3526 8532	4940 0302	L		_
L	8		3 8	653		976	1323	1702	1818		7037			3760		4281			000	<i>7</i> E	- C	,, ,	~ 1 5	e: 19	4120	-		- 12	1385	اذ

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Lynny sanians chromosome 21 segment HS210084	Domo seriens chamosome 21 segment NS21C082	Homo septiens chromosome 21 segment HS21C085	Homo septems throughout protein (AF038169), mRNA	EST68385 Infant brain Homo sapiens cDNA 5 and similar to similar to heat shock protein, so Note	hingage x1 NCI CGAP_Lu24 Home sapiens cDNA done IMAGE:3168120 3' similar to contains minimary	MER18 repetitive element : https://doi.org/10.1016/10.0016	MECHANIST STATE OF THE STATE OF	PLANCAGE ANCI COAP LIZA Homo sepiens CDNA clone IMAGE:3166120 3' similar to contains MEKING.	Interessing Comment:  E4-74 at Some helts NHT Home sectors CONA done IMAGE:757422 5	Lymp sanions eukarvatic initiation factor 4AI (EIF4A1) gene, partial cds	DAILE STATE OF THE COLORFE (KIAA0933)	Hums sanlens chromosome 21 segment HS21C084	DKF2076181710_r1 761 (synonym: hamy2) Homo sapiens CDNA done DKF2p761B1710 o	258 1 KDA PROTEIN C21ORF6 (KIAA0933)	RAMIN SPECIFIC ANGIOGENESIS INHIBITOR 2 PRECURSOR	Home sapiens CCAAT-box-binding transcription factor (CBF2) mRNA	Homo sapiens CCAAT-box-binding transcription factor (CBF2) miNNA	qy64e11.x1 NCI_CGAP_Bm25 Home saplens cDNA clone IMAGE:2010612.9 suring	MER12 repetitive element:  Homo seniens basic transcription factor 2 p44 (bt/2p44) gene, partial cds, neuronal apoptosis inhibitory	protein (naip) and survival motor neuron protein (srnn) genes, complete cas	Home saplens nucleonar prospring with the complete ods	Homo saplens nucleolar phosphophore B22 (14) 111 April 114 April 1	601058602F1 NIH MGC 10 Homo septems Control Control (1974) MGE 4414029 3' similar to conteins	2135503.s1 Soares pinesi gland N3HPG Home segrens control control	Alu repetitive element contains element mentat la produce de la la la la la la la la la la la la la		we78h03.x1 Scenes_Dicotograefe_colon_NHCD Homo septiens curva curva innoverses_Dicotograefe_colon_NHCD Homo septiens element PTR6 repetitive element; SW:RL29_HUMAN P47914 608 RIBOSOMAL PROTEIN L29 ;contains element PTR6 repetitive element;	
	Top Hit Databerse Source					NAM US IMAN	Т	EST_HUMAN	EST HUMAN	SWISSPROI	EST HUMAN	EST HUMAN		SWISSPRO	T LINGS	ESI DOMON	SWISSPROT	SWISSING			EST HUMAN				EST HUMAN		EST HUMAN	ESI TIOMAN	EST HUMAN	
<u> </u>		_			ž		3	EST	ES	3	ES			<u></u>			5				쁴	Z	본	본	Ü	F	-	4		1
CAL BIRTH	Top Hit Acession No.				163285.2	N 81/8636	4350878.1	222239.1	-222239.1	23249	E22238.1	A442272.1	F175325.1	SOYSRE	AL 163284.2	4L118573.1	Q9Y3R5	D60241 SW	5031824 NT	20100	Al356086.1	U80017.1	M28699.1	M28699.1	BE535440 1		AA719297.1	AW867740.1	A1870074 4	Tan Jan Jan Jan Jan Jan Jan Jan Jan Jan J
+	Most Similar (Top) Hit To BLAST E	Aging	5.0E-09 AL163284.2	4.0E-09 AL163282.2	4.0E-09 AL163285.2	4.0E-09	4.0E-09 A	3.0E-09 BE222239.1	3.0E-09 BE222239.1	3.0E-09 P23249	3.0E-09B		۹.	$\sim$			2.0E-09 (		1.05.09	1.05-09	1.0E-09	1.0E-09	4 OF-09	4 PE-09	90 30	1.05-09	1.0E-09	9.0E-10	6	8.0E-10
	Mos Expression (1 Signal BI		1.29	1.41	2.28	2.05	17.61	3.39	1.04	0.92	1.29	6.3	4.42	1.63	13.77	14.91	8.68	3.1	1.68	1.68	1.35	9,	1 22	17.33		0.08	6.25	1.94		7.26
	ORF SEQ E		11899	-		11495	12463	12383	12563	12855	13289		14281	14351	11280		12359		11128	11129		_				12998		11333		12806
	SEQ ID		8808	5552	2968	8438	7336	7265	7450	7541	7908	8303	1000	8372		L	L		L	609	7389	_	١	l		3 7984	0640			7781
	Probe SEQ ID 8		1848	247	920	144	2362	2280	2482	2578	300	2002	2027	4381	1239	4848	2265	3834	1092	1092	2428		2819	2855	2855	2986	7987	1280		2760

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Table 4
Single Exon Probes Expressed in HBL100 Cells

														_				<u> </u>			$T_{\ell}$	1	۴	_	1 1	7	H	۳	H	5	6	1
	Top Hit Descriptor	HOWAY MCMA) and DNA-PKcs (PRKDC) genes, partial cds	Figure Edition in Maria Sapiens CDNA 6 and	Homo segiens TPA Inducible protein (LOC51586), mRNA	Homo seplens TPA inducible protein (LOC51588), mRNA	LYSP100 PROTEIN (LYMPHOID-RESTRICTED HOMOLOG OF SP100)	LINE-1 REVERSE TRANSCRIPTASE HOMOLOG	H. saplens DHFR gene, excn 3	Homo septens ASCL3 gene, CEGP1 gene, C11or114 gene, C11or115 gene,	gens #02407.x1 NCI CGAP Pr28 Homo septens cDNA clone IMAGE:2086021 3*	RC3-CT0254-031059-012-912 CT0254 Homo septens cDNA	DKEZ0434N219 11 434 (synonym: htes3) Hamo septens cDNA clane DKFZP434N219 o	HYPOTHETICAL GENE 48 PROTEIN	Home seciens WRN (WRN) gene, complete cds	Another of Search a Blackweeks 2NbHP8tc8W Home septems on A done IMAGE:1758CAP 3	Notice to contains LTR8.b2 LTR8 repetitive element; series to contains LTR8.b2 LTR8 repetitive element; series to contains LTR8.b2 LTR8 repetitive element;	HIGHER AND COAP GCS Home sapiens cDNA clone IMAGE:2949844 3' similar to contains Att	repetitive element	Homo sapiens chromosome 21 southern contains china done IMAGE:272963 3' similar to contains	yy32f06.e1 Soares melanocyte ZNDHM Florino explores control and a second	-	_	MAJOR CENTROMERE AUTOANTIGEN B (CENTROMERE PROTEIN OF CENTROMERE PROTEIN OF CENTROMERE INHBITION	Homo sapiens basic transcription factor 2 p44 (bt/2p44) gene, perual cus, neulonia appropria	protein (neip) and survival motor neuron protein (smin) genes, compress con- leo2138640F1 NIH_MGC_83 Home sapiens cDNA clone IMAGE:4273377 67	MR0-SN0038-290300-001-f01 SN0038 Home saplens CDNA	AV652123 GLC Homo saplens cDNA clare GLCCAN I I S	QV0-CT0225-191199-058-e08 CT0225 Home Sapiens CDNA	QV2-T10003-161189-013-910 T10003 Hamo septems CDNA clare DKF2p434N1317 6			
and a row a significant	Top Hit Database Source		L	ESI HUMAN	_	CWISSPROT	SWISSPROT	12		NA FOR	EST CINAN	MALAN TOTAL	ESI HOMAN	SWISSPROI	ž	EST_HUMAN	EST HUMAN	EST_HUMAN	N		EST HUMAN	SWISSPROT	EWISSPROT		TN FOR	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	_
AT BIRING	Top Hit Acession No.		1	AA376832.1		7/00//	T	7.0E-10 P08547			6.0E-10 AI424405.1	1	5.0E-10 AL046804.1	T		4.0E-10 AI221083.1	0 AA515260.1	0 AW 594709.1	4 OF-10 AL 163303.2			3.0E-10 AY005150.1	10 148800	101748988	10 U80017.1	2.0E-10 BF675047.1						
	Most Similar (Top) Hit BLAST E	Value	8.0E-10	8.0E-10	7.0E-10	7.0E-10	7.0E-10	7.0E-10	7.0E-10	6.0E-10	8.0E-10	6.0E-10	5.0E-1(	5.0E-1	5.0E-1		4.0E-10 AA						_1				1		3 1.05-10		1.0E-10	
	Expression Signal		11.08	1.87	39.41	39.41	1.69	9.17	2.59	2.89	1.11	2.52	4.91	1.15	1.25	1.48		4.48			1.72			1.49	2.98	1.37					0.97	1.30
	ORF SEQ ID NO:		10228	14059	10732	10733	11643		13041	10956	12683			13429			10607			7 12582	10958	L	10102	6 10103	<u> </u>	9,		57 11629		28 13454	8470	8470
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Page 70 of 209 Table 4 Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Home septens ruichear factor of kappa light polypapides gene entrances in Brossa (VIII 1907), secret	eds (DUSP9), ribosomal specificity phosphetase 9 (DUSP9), ribosomal	Homo septems X28 region near ALL Notes Community protein kinase I (CAMICI), creatine transporter (CRTR), protein L18a (RPL18a), Ca2+/Calmodulin-dependent protein kinase I (CAMICI), creatine transporter (CRTR), contain (CDMI), advanced today speciel >	Committee of Discontinuous Al Discontinuous containing dual specificity phosphatase 9 (DUSP9), ribosomal	From separate Act region (22+/Celmodulin-dependent protein kinase I (CAMKI), creatine transporter (CRTR), protein L18s (RPL18a), Ca2+/Celmodulin-dependent protein kinase I (CAMKI), creatine transporter (CRTR),	CDM protein (CDM), adrendeukodystrophy protein 2  CDM protein (CDM), adrendeukodystrophy protein 2  CDM protein (CDM), adrendeukodystrophy protein 2  CDM protein (CDM), adrendeukodystrophy protein 2	Home saperities of the second	11 2-HT0203-291099-018-008 HT0203 Homo sapiens cDNA	DIKEZAZZOSS 1 647 (synonym: hfbrt) Homo sapiens cDNA clone DKFZp547DZZ5 6	DNR F2647D225 r1 647 (synonym: htbr1) Homo sapiens cDNA done DKF2p547D225 6	DIXE 70:5470225 1 547 (synonym: hfbr1) Hono saplens cDNA clone DKFZp5470226 6	DIX F77.5470225 11 647 (synonym: hfbrt) Home saplens cDNA clone DKFZp5470225 6	oo7aff/1 st Strategene schizo brain S11 Homo saplens cDNA clone IMAGE:97029/ 3	NA5311.61 Soares adult brain N2b5HB55Y Homo septens cDNA clone IMAGE:172173 3' stmiter to contents	1.1 repetitive element :	FST34392 Embryo, 6 week I Homo saplans cDNA 5' end	Human matrix Gla protein (MGP) gene, complete cds	Human matrix Gla protein (MGP) gene, complete cds	Homo saplens chromosome 21 segment HS21C083	Hano saplens chromosome 21 segment HS21 C063	ALDEHYDE OXIDASE	Zu01b12.r1 Soares_bestis_NHT Homo sapiens Corty orders in 1200 1205 5	601607631F1 NIH_MGC_/1 Hario september 2017	Homo sapiens chromosome 21 segment M3210M4/	HUMSUPY069 Human brain cDNA Homo sapiens CONA COLO 2000 HUMANA HU	Mus musculus expressed in non-metastatic deals 4, protein (Antaco) (Missing expressed in non-metastatic deals (Missing expressed in non-metastatic deals (Missing expressed in non-metastatic deals (Missing expressed in non-metastatic deals (Missing expressed in non-metastatic deals (Missing expressed in non-metastatic deals (Missing expressed in non-metastatic deals (Missing expressed in non-metastatic deals (Missing expressed in non-metastatic deals (Missing expressed in non-metastatic deals (Missing expressed in non-metastatic deal	W35406X1 NCI_CGAP_Kid12 Hamo sapiens conva cidire invacionalia	Г	Г	1	
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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	yg43e12.r.1 Sogres Infant brain 1NIB Homo septems cDNA clone IMAGE:35144 6	19439-12 of Sogres Infant briefin 1NIB Home espheres count done invocation and offerbox receptor-like protein	Gallus galus moglobih, beta-h grozin, beta-h grozin, epsivar grozin, and allos galus moglobih, beta-h grozin, complete cds COR3'beta (COR3'beta) genes, complete cds	Gallus gallus rho-globin, beta-H globin, beta-A globin, epsilon-grobin, and chastury receptur nine process. COR3 beta (COR3 beta) genes, complete cds	Human endogenous retrovints HERV-P-1470	RETROVIRUS-RELATED GAG POLITPROTEIN (VENSION 1)	THE SCORE AT LINE IN THE STREET OF THE STREE	Homo sapiens FRA3B common fragile region, diadenosine triphosphate hydrolase (FHIT) gene, exon 5	KC3-B10316-170200-014-803 B103101 initial organical control of the	Home septens difference at a barose Lower equipme china	QV2-B10256-201089-414-801 B10230 multip septiens control	Hamo sagrans out, garle notes	Homo sapiens crudinosome z 1 seginan 102 roco	Home separate DROADTR mRNA, complete cds	DAVETERNI RINDING PROTEIN	UANDA caniens homonantisate 1.2-dioxygenase gene, complete cds	CAIN BRIGHTS 170300-202-412 BN0105 Homo suplens cDNA	A KIN SPICI II F MATRIX PROTEIN PRECURSOR (LSM34)	34 NO STATE HOME Septens CDNA clone HTFAW F06 6	1728111,51 NCI CGAP GCB1 Homo sapiens cDNA clone IMAGE:1302573 3' similar to contains Alu	repetitive element	Human chromosome 21 distal rang 4mil DIVA	EST04462 Feta brain, Stratageria (carrespond) Total Appe IMAGE 2201217 6	E42505.y1 NCI CCAP Binaz natio septens control and the septens of	Homo sapiens Aq pseudosumsumsumsumsumsumsumsumsumsumsumsumsums	2/4911.St Sources jetted liver enlacen 1NEIS S1 Home septients CDNA clone IMAGE:460676 3	2/4911.81 SOLICE JULY Homo septem CDNA clore IMAGE:2270745 3' similar to TR.013639 Q13639	MARINER TRANSPOSASE.;
	Top Hit Database Source	HUMAN	EST_HUMAN				SWISSPROT	EST_HUMAN		EST HUMAN		EST HUMAN				10000	SWISSPRO	1444	EST FIONAN	SWISSPROI		EST HUMAN		EST HUMAN	EST HUMAN	1	EST HUMAN	EST HUMAN	EST HUMAN
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	Expression Signed	3.84	3.84	3.97	902	0.83	5.68	0.74	0.94	1.04	0.76	1.84	67.0	1.24	2.94	1.4	1.14	2.2	0.93		0.75	9.87						6.02	0.7
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historio x1 NC\_CGAP\_GU1 Homo sapiens cDNA clone IMAGE.2970040 3' similar to contains MER18.t1 Homo sapiens X-linked anhidrotic ectodermal dysplasta protein gene (EDA), exon 2 and flanking repeat Indiadol x1 Soures\_NFL\_T\_GBC\_S1 Homo septens cDNA clone IMAGE:2809377 3' similar to TR:014517 014517 SMRP: hd13d01.x1 Soures\_NFL\_T\_GBC\_S1 Hamp suplens cDNA clone IMAGE;2808377 3' similer to TR:014517 wm51f07.x1 NCI\_CGAP\_U/2 Homo septems cDNA clone IMAGE:2438493 3' similar to contains L1.b3 L1 Homo saplens basic transcription factor 2 p44 (btf2p44) gene, partial cds, neuronal apoptosis inhibitory 277712.81 Seeres testis\_NHT Homo saplens cDNA clone IMAGE:728350 3' similar to contains Atu y82/104.11 Soares placenta Nb2HP Homo saplens cDNA clone IMAGE:145769 6 Homo sapiens glypican 3 (GPC3) gene, partiel cds and flanking repeat regions x48407.r1 Soares\_testis\_NHT Homo sapiens cDNA clone IMAGE:795469 5 Homo saplens testa epecific Testis Transcript Y 2 (TTY2) mRNA, partial cds Homo saplens testis-specific Testis Transcript Y 2 (TTY2) mRNA, partial cds AD132248 NT2RP3 Homo saplens cDNA clone NT2RP3004070 5 protein (naip) and survival motor neuron protein (smn) genes, complete ods Homo sapiens chromosome 21 segment HS21C007 Homo sepiens CST gene for cerebroside sulfotransferase, exon 1, 2, 3, 4, 6 AU132248 NT2RP3 Homo saplens cDNA clone NT2RP3004070 5 repetitive element, contains element MER22 repetitive element; PM2-HT0224-221099-001-e11 HT0224 Homo sepiens cDNA Mus musculus keratin-essociated protein 6.2 (Krtap6.2), mRNA Top Hit Descriptor Rat U3A small nuclear RNA CM0-BT0281-031199-087-403 BT0281 Homo sepiens cDNA |LE-UN0071-120400-085-605 UN0071 Homo septens cDNA Homo saplens Xq pseudoautosomal region; segment 1/2 Homo saplens prion protein (PrP) gene, complete ods Homo sapiens prion protein (PrP) gene, complete ods TBX15 PROTEIN (T-BOX PROTEIN 15) Single Exon Probes Expressed in HBL100 Cells MER 18 repetitive element; Rat U3A small nuclear RNA 014517 SMRP. regions EST HUMAN EST HUMAN EST\_HUMAN EST HUMAN EST\_HUMAN EST HUMAN EST HUMAN EST\_HUMAN **FUNAN** SWISSPROT SWISSPROT EST HUMAN EST HUMAN Top Hit Database Source Ę Z 눈 Ę 6754495 NT 4.0E-13 AW378614.1 3.0E-13 AF003528.1 Top Hit Acession 4.0E-13 AF003529.1 4.0E-13 AA454054.1 5.0E-13 AA435773.1 6.0E-13 AL163207.2 6.0E-13 R78338.1 1.0E-12|AW627874.1 1.0E-12 AU132248.1 1.0E-12 AU132248.1 AB028900.1 2.0E-12 AW802131.1 2.0E-12 6754 1.0E-12 AF000991.1 1.0E-12 AF000991.1 9.0E-13 AJ271735. 3.0E-12 AW341683.1 3.0E-12 AW341683.1 1.0E-12 AIB71728.1 8.0E-13 U80017.1 BE063509.1 8.0E-13 U29185.1 U29185. ģ 2.0E-12 J01884.1 2.0E-12 J01884.1 2.0E-12|070306 070306 8.0E-13 2.0E-12 BLAST E Value 五多)压 1.02 4.21 236 3.41 0.85 0.67 29.33 4.84 1.34 1.53 4. Ž 1.65 1.34 1.12 3.43 0.75 0.85 0.85 2.34 3.43 Expression Signal 11883 10747 12137 13789 13028 13854 10205 13027 14712 14713 11673 13418 10641 ORF SEQ ID NO: 5242 7362 9592 6820 8324 8261 8261 8845 8788 8015 8786 8015 8550 6192 6927 8994 9727 9289 8994 9727 6839 5639 6609 8394 SEQ ID 179 3313 2391 1802 3248 3543 3843 3783 20 707 2897 2897 4742 4297 4742 133 922 3386 3998 3998 612 SEQ ID ÿ

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	Top Hit Descriptor	2468g08.r1 Soares lestes Juni Praint 2/2	Homo sapiens At process 21 segment HS21C010	Homo saplens chromoson in 27 100 Homo saplens cDNA	CM3-F L0100-1400-20-20-20-20-20-20-20-20-20-20-20-20-2	ob18402.61 NCI CGAP, Not Turned A Dipous containing dual specificity phosphatase 9 (DUSP9), riposantial	Homo sepiens XXS region from Annual Through the Annual (CAMIN), Greating Language (CAMIN), Greating (C	protein L188 (Kr. L199), carendeutodystrophy protein > CDM protein (CDM), adrendeutodystrophy protein >	Danjo rerio fibroblast grown racka recent racka recent racka recent rece	ONA polymerase delta small subunit (POLD2) gene, excris 1 triougn 11 and contraction of the contraction of t	nab76/05.xt Source NSF F8 9W OT PA P S1 Home septents CLYN curror in the profession of the part of the profession of the part of the profession of the part of the profession of the part of the profession of the part of the profession of the part of the profession of the part of the	Homo saplens S164 gene, partel cus, roll mis light	partial des	Homo saplens chromosome 21 sources. Homos, Genomic, 342 nt, segment 2 of 2]	FGF-1 Florolius ty Commission 18 and 14	Homo saplens LGMUZE 1942 1 IPP2 LMP2, TAP1, LMP7, TAP2, DOB, DQB2 and NINCS, 5, 12	H.saprens Divina, Divi	+	A PARAMSBOOGE NO. CGAP. Brn64 Homo sapiens cDNA clone IMANCE: 4 100000	7		T	_	П	_	П	Seguinus coultage in 24 Home septens cDNA clone IMAGE 21 1271 1271 1271 1271 1271 1271 1271		- 1		•
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	Top Hit Descriptor	x677810.x1 NCL_CGAP_Gas4 Homo septens cDNA clone IMAGE:2823148 3' similar to contains MER10.t2	MER10 repotitive element;	Homo sapiens FRA3B common fragile region, diadenosine triphosphate hydrolese (FHI I) gene, exprise Homo sapiens FRA3B common fragile region, diadenosine triphosphate 1 (MI II TIDRUG RESISTANCE.	CANALICULAR MULTISPECIFIC ORGANIC ANION I NANSPONTENT (MICHAEL PROTEIN)	ASSOCIATED PROTEIN 2) (CANALICOLDAN MICE) MAGE: 2575185 3' similar to contains L1.02 L1	repetitive element;	Homo sapiens LGMD2B gene	2/67 e.06.r1 Soares_pregnant_ulanus_NbHPU Homo sapaens CUNA clune IMAGE-279190 3' samilar to	W/73c12.61 Soares_multiple_sciencels_ZNbHMSP Homo sapens con constitution	contains L1.13 L1 repetitive element ;	Home saplens Xq pseudoautosomal region; segment 2/2	Homo sapiens Xq pseudoautosomal region; segment 2/2	Homo sapiens chromosome 21 segment HS21C103	RC5-BT0377-091299-031-D12 BT0377 Hamp sapients CATOR1), mRNA	Homo sapiens mabdoid tumor deligibation to a contraction of the contra	Homo sapiens chromosome 21 segment in 2210046	Homo sapiens chromosome 21 segment H221088	Homo sapiens chromosome 2.1 eaginement HS21CD68	Homo sapiens chromosome 21 segundan 122.	Homo saplens chromosome A region in the company of	(G6PD) gene, compress cas s Homo septens chromosome 21 segment HS21C103	Home septens ribosomal protein L23A (RPL23A) gene, compression	Т	Γ	Г	Г	Г	П	
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t		<u> </u>	7.0E-14 AW		1.0E-14/	5.0E-14 Q63120	5.0E-14	4.0E-14 P04928	4.0E-14	4.0E-14	4.0E-14	3.0E-14 X	2.0E-14	2.0E-14	2.0E-14	4.VE	20E-14	44 10 1	100	1.05.14	2	1.0E-14	1.0E-14	1.0E-14	1.0E-14	1.01-14	1.0	1.08-14	1.0E-14	47
	Most Similar (Top) Hit BLAST E	Value	_									2				* 1	<u></u>		2 2	5 2	5	8	4.33	18	7	6.58	9.58	듸	81	
	Expression Signal		2.98		13.88	4.27	1.45	1.59	6.27	0.88	0.8	1.12	2.33	2.33		1.24						21.63								
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			+-	1	6416	5641	1 18	7739	6828	8671	8	26.5	5427	5427	77.27	7298	7366	7425	6909	6379	6379	2041	140		88	8110	8110	8795	8339	6238
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	Top Hit Descriptor	Homo septiens transcription factor IGHM enhancer 3, JM11 protein, JM4 protein, JM5 protein, 134	JM10 protein, A4 differentiation-dependent protein, triple Linn derivain process. Complete cds; and L-type calclum channel 42-	80114863ZF1 NIH MGC 18 Homo sapiens curving and an income of the control of the c	Homo saplens Xq pseudoeutosomai regirar, segment	Homo saplens chromosome 21 segment no 21000	herma 2A-like protein gene, hereditary hasmochromatosis	Human hereditary haemochromatosis regent, instantial trasporter (NPT3) gene, complete cds (HLA-H) gene, RoRet gene, and a sodkim physicitate trasporter (NPT3) gene, complete cds (HLA-H) gene, RoRet gene, and Roshing HS310103	Home eaplens chromosome & eagurer.	LY1142F FULLIA IOCT.	NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5 NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5 NAME   NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 5	Homo sapiens calcium channel alpha 1E subunit (CACATA I E) gara,	spliced Homo saplens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively	spliced Home septens calclum channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively	spliced https://www.nci.cgap_Kid13 Homo sapiens cDNA clone IMAGE:3146258 3' similar to contains MER29.b3	_	$\neg$	spliced Homo sepiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, atternatively Homo sepiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, atternatively	spiloed spiloed NCI CGAP_HN10 Homo septens cDNA clone IMAGE:2741521 3' similar to contains L1.t3 L1			N MARINER TRANSPOSASE N MARINER TRANSPOSASE N ht40e02.y1 NCI_CGAP_0v34 Homo septems cDNA clone IMAGE:2899162 6*	l
	Top Hit Detabase Source	_	<u> </u>	EST HUMAN	Z	5		IN.	Ę		SWISSPROT	_	Į,	Ę	Ę	EST HUMAN	EST HUMAN	Ę	Ę	EST HUMAN	EST HUMAN	EST HUMAN	
}	Top Hit Acession No.		10220	807 (8.1	Ī	T	02500.5	328.1	63303.2				2.0E-15 AF223391.1	2.0E-15 AF223391.1	AF223391.1	2.0E-15 BE350127.1	BE350127.1	5 AF223391.1	5 AF223391.1	5 AW 238499.1	6 AI806335.1	15 AI689984.1	la peropera
	<b>⊉</b> + ₪	Agine	i i	9.0E-10 AF	0.0C-13 E	6.05-10	5.0E-13 AL	A 0F-15	4.0E-15		3.0E-15	3.0E-15 P92485	2.0E-15	2.0E-15	2.0E-15 A	2.0E-15	2.0E-15 B	2.0E-16	2.0E-15	2.0E-15	5 2.0E-15		1.05-15
	Signal E		•	20.	1.02	4.97	4.86		243		12	2.28	2.86	2.77	2.77	1.11	1.11	0.91	0.91		. 255	1.94	1.78
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	0	j Ž		7088	5513	5994	5444		48	2087	0606	9754	6308	5448	5416	l		_	1	<b>I</b>		<b>\</b>	Ш
	- 0	 Ö	-	2108	2739	926	407		2688	424	4096	4770	2		307	2244		3	3450	3430	3804	2002	2843

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Mus musculus offactory receptor cluster, OR37A, OR37B, OR37C, OR37E genes and OR37D pseudogene or offactory receptor cluster, OR37B, OR37B, OR37B, Selection of or offactors o Homo sapiens TSX (TSX) pseudogene, exon 6 ZONADHESIN PRECURSOR ENVELOPE GLYCOPROTEIN GP340 (MEMBRANE ANTIGEN) (MA) [CONTAINS: GLYCOPROTEIN Homo sapiens FRA3B common fragile region, diadenosine triphosphate hydrolase (FHIT) gene, exon 5 AV681393 GLC Homo sapiens cDNA clone GLCGSA01 3' af39g11.s1 Soares\_total\_fetus\_Nb2HF8\_9w Homo saplens cDNA clone IMAGE:1034084 3' stmiler to df45c01.y1 Morton Fetal Cochlea Homo sapiens cDNA clone IMAGE:2486376 6' DKFZp434P037\_11 434 (synonym: hiss3) Homo sapiens cDNA clone DKFZp434P037 5' Homo sapiens pituitary tumor transforming gene protein (PTTG) gene, complete ods EST06060 Infert Brain, Bento Sceres Homo septens cDNA clone HIBBA13 6' end PM4-BT0650-010400-002-509 BT0550 Hamo sepiens cDNA df45c01.y1 Marton Fetal Cochlea Hamo sepiens cDNA clone IMAGE:2486376 6 RC3-HT0649-100500-022-b05 HT0649 Hamo sepiens cDNA wr86e04 x1 NCI\_CGAP\_Kid11 Hamo sepiens cDNA clans INAGE:2494590 3' EST384702 MAGE resequences, MAGL Hamo sepiens cDNA Homo sapiens gens for TMEM1 and PWP2,complete and partial cdt QV1-UM0036-200300-116-902 UM0038 Homo sepiens GDNA QV1-UM0036-200300-116-902 UM0038 Homo sepiens GDNA MYELIN-OLIGODENDROCYTE GLYCOPROTEIN PRECURSOR QV0-BN0148-070700-283-s10 BN0148 Homo saplens GDNA CM1-NN1003-200300-153-e01 NN1003 Homo saplens GDNA QV0-OT0032-080300-165-d01 OT0032 Homo saplens GDNA Human SSAV-related endogenous retroviral LTR-like element Top Hit Descriptor PM4-BT0650-010400-002-909 BT0650 Homo saplens cDNA H.saplens DNA for endogenous retroviral like element Homo saplens chromosome 21 segment HS21C079 LINE-1 REVERSE TRANSCRIPTASE HOMOLOG contains OFR.12 OFR repetitive element. contains element L1 repetitive element; Single Exon Probes Expressed in HBL100 Cells EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST\_HUMAN EST\_HUMAN SWISSPROT EST\_HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST HUMAN EST\_HUMAN EST\_HUMAN SWISSPROT SWISSPROT EST HUMAN SWISSPROT Top Hit Detabase 눌 눋 1.0E-16 AA628592.1 1.0E-16 BF327942.1 8.0E-17 AW880701.1 Top Hit Acession 9.0E-17 AW900048.1 2.0E-16 AL163279.2 2.0E-16 J03061.1 AF200719.1 3.0E-16 AF020503.1 4.0E-16 AW797168.1 4.0E-16 Q16653 3.0E-16 AV681393. 3.0E-16 AW022862. AF135446.1 2.0E-16 X89211.1 1.0E-16 AF200719 4.0E-16 AW797168.1 3.0E-16 AL046445.1 1.0E-15 P08647 1.0E-16 BE182898.1 1.0E-16 Al984928.1 6.0E-16 AW972911.1 AW972611.1 6.0E-18 AJ251154.1 6.0E-16 AAB92176.1 4.0E-16 AB001523.1 3.0E-16 P03200 3.0E-16 T08169.1 3.0E-16 4.0E-16 (Top) Hit BLAST E Value 25.58 **1.8**3 0.89 1.7 0.78 1.08 2.54 423 0.67 4.76 3.79 2.13 0.97 1.58 0.98 Ē 0.83 0.83 6.06 5 23 Expression 12009 13665 13839 14753 14036 14003 12413 13403 14002 10215 12885 12412 11617 13099 14212 14917 ORF SEQ Ö N Q 6913 8660 6012 5456 8863 5973 9048 5247 7926 9769 7579 8832 5500 6423 9015 5199 8383 9015 7292 7292 6458 7 8085 9228 9940 7061 SEO ID 3654 1927 2617 3851 4785 184 38307 957 1426 4019 133 133 463 3375 4019 23178 3069 4063 2080 2080 <u>₹</u> SEO ID ÿ

Page 77 of 209 Table 4 Probes Expressed in HBL100 Cells

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Single Exon Probes Expressed in Tibracy Cons	Top Hit Descriptor	Homo sepiens chromosome 21 segment HS21C080	Miss misserilus excliposinolein B editing complex 2 (Apobeo2), mRNA	DOL HINDRY 22000 021-004 HINDOO3 Homo saplens cDNA	Action of Statement Inna (#037210) Hamp sapiens cDNA clone IMAGE:78839 5	younder it at all the post of the same same of the IMAGE:1058528 3'	nighedos si Noj Coar Colo Indio agricio Con Cone IMAGE:2804784 3'	XGOSGUBAXI SORIES IN L. C. C. C. C. C. C. C. C. C. C. C. C. C.	MASSACE LED STINCTER COMP. Septens cONA clone IMAGE:3181999 3'	HENDEROA Y INC. CGAP 1.1/24 Homo saplens cDNA clone IMAGE:3181999 3'	Anterson St. NCI CGAP Eso2 Homo sapiens cDNA clone IMAGE: 1959922 3' similar to contains Alu	repetitive element	qt63a08.x1 NCI_CGAP_Esoz Homo sapients cultin dulle innoce.	repetitive element.	MOINTEN DOMESTING THE COMPANY AND THE COMPANY		CUNALITIES IN THE STATE OF THE PROTEIN (200 KDA NEUROFILAMENT PROTEIN) (NEUROFILAMENT	HEAVY POLYPEPTIDE) (NF-H)	Homo septens X28 region neer ALD locus containing dual specificity prospirations 9 (Coor of, second	protein L18a (RPL18a), Ca2+/Caimodulin-dependent protein kinase I (CAMKI), creatine transporter (L7.17), CDM protein (CDM), adrendeukodystrophy protein (CDM), adrendeukodystrophy protein (CDM), adrendeukodystrophy protein (CDM), control (CDM), adrendeukodystrophy protein (CDM), control (CDM), adrendeukodystrophy protein (CDM), control (CDM), adrendeukodystrophy protein (CDM), control (CDM), co	MULTIDRUG RESISTANCE PROTEIN (CELLOCITED)	Homo sapiens Xq pseudoautosomal region; segment 2/2	Homo sapiens chromosome 21 segment nozirovo	COLLAGEN ALPHA 1(III) CHAIN PRECONSON	Homo saplens thrombospondin 2 (1 HBS2) gene, promoter region and the conjugating enzyme E2D 3	Homo sapiens mannosidase, beta A, tysosoma (MANDA) gens, and unquini congress.	(UBE2D3) genes, complete cds V20c07 rt Scares febril liver spleen 1NPLS Homo septens cDNA clone IMAGE:128389 6	Homo septems protein tyrosine phosphatase, non-receptor type substrate 1 (PTPNS1) mRNA	xc10b04x1 NCI COAP Pant Homo sapiens cDNA clone IMAGE:2837071 3' similar to gb:1.20808 dus	RIBOSOMAL PROTEIN L4 (HUMAN);  ACIDICAL XI NCI CGAP Part Homo septens cDNA clone IMAGE:28370713' similar to gb:1.20868 60S	
- FIGDES E	Top Hit Detrabase Source				ESI HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	SWISSPROI	HOLD HOMAIN	ESI TOWAN	EST HUMAN		EST HUMAN	EST HUMAN	SWISSPRO	SWISSPROT	SWISSPROT		<b>z</b>	SWISSPROT	NT	NT	SWISSPROT	N.		FA FOR	TOWN THE	Ž -	EST_HUMAN	EST_HUMAN
Single Exc	Top Hit Acession No.	1469280 2 NT	1004004	ğ	1			3.0E-17 AW119123.1 E		3.0E-17 BE326522.1	1	1270080.1			A722832.1	128983	128983	P12036  S		U52111.2	P08183	AJ271738.1	AL163207.2	P02461	U79410.1		AF224669.1	R09942.1	4768977 N	AW316978.1	AW316976.1
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	Expression Signed	+	2.80	1.93	5.89	2.71	48.0	1.06	1.31	1.36	1.36	2.62		2.83	2.23	1.92	1.92	278	3	4.97	000	3.03	4 28	4 54	90.0	2.00	1.05	8.44	1.84	65.43	65.43
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Page 78 of 209 Table 4 obes Expressed in HBL100 Cells

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Single Exon Probes Expressed in the control of the	Top Hit Descriptor	Ratus novedicus partal Convincio (Transferase (TISSUE TRANSGLUTAMINASE)	(TGASE C) (TGC)	qm65g11xd Sogres, placenta, globweeks, ZNBHFowery Franco	similar to contains Au repoured complete a DNA clone IMAGE:3030611 3' similar to contains MED Au MAGE:3030611 3' similar to contains MED Au MAGE:30 MA	MERZ9 repeature element. ho36h04x1 MCI_CGAP_Utt Homo septens cDNA clone IMAGE:3039511 3' similar to contacts millioned to contacts and contacts and contacts and contacts and contacts and contacts are contacts and contacts and contacts are contacts and contacts are contacts and contacts and contacts are contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and contacts are contacts and	MERZO repetitive escribara. nq24f11.s1 NCI_CGAP_Coto Homo septens cDNA clone IMAGE:1144845 3' similar to galantees.	KERATIN, TYPE I CYTOSKELE I AL 19 (1907)  KERATIN, TYPE I CYTOSKELE I AL 19 (1907)  N-ACETYLACTOSAMINIDE BETA-1, BARANCHING ENZYME) (1907)	ACETYLGLUCOSAMINYLI KANSTERVELY (CETTYLGLUCOSAMINYLTRANSFERASE (N-N-ACETYLACTOSAMINIDE BETA-1, ACETYLGLUCOSAMINYLTRANSFERASE (N-ACETYLACTOSAMINIDE BETA-1, ACETYLGLUCOSAMINYLTRANSFERASE (N-ACETYLACTOSAMINING) (GNT)	ACETYLGLUCOSAMINYLTKANS/FEXXXII) (17.7) ACETYLGLUCOSAMINYLTKANS/FEXXIII) ACETYLGLUCOSAMINYLTKANS/FEXXIII) ACETYLGLUCOSAMINYLTKANS/FEXXIII) ACETYLGLUCOSAMINYLTKANS/FEXXIII) ACETYLGLUCOSAMINYLTKANS/FEXXIII) ACETYLGLUCOSAMINYLTKANS/FEXXIII) ACETYLGLUCOSAMINYLTKANS/FEXXIII) ACETYLGLUCOSAMINYLTKANS/FEXXIII) ACETYLGLUCOSAMINYLTKANS/FEXXIII) ACETYLGLUCOSAMINYLTKANS/FEXXIII) ACETYLGLUCOSAMINYLTKANS/FEXXIII) ACETYLGLUCOSAMINYLTKANS/FEXXIII) ACETYLGLUCOSAMINYLTKANS/FEXXIII) ACETYLGLUCOSAMINYLTKANS/FEXXIII) ACETYLGLUCOSAMINYLTKANS/FEXXIII) ACETYLGLUCOSAMINYLTKANS/FEXXIII) ACETYLGLUCOSAMINYLTKANS/FEXXIIII) ACETYLGLUCOSAMINYLTKANS/FEXXIIII ACETYLGLUCOSAMINYLTKANS/FEXXIIII ACETYLGLUCOSAMINYLTKANS/FEXXIII ACETYLGLUCOSAMINYLTANS/FEXXIII ACETYLGLUCOSAMINYLTANS/FEXXIII ACETYLGLUCOSAMINYLTANS/FEXXIII ACETYLGLUCOSAMINYLTANS/FEXXIII ACETYLGLUCOSAMINYLTANS/FEXXIII ACETYLGUCOSAMINYLTANS/FEXXIII ACETYLGUCOSAMINYLTANS/FEXXIII ACETYLGUCOSAMINYLTANS/FEXXIII ACETYLGUCOSAMINYLTANS/FEXXIII ACETYLGUCOSAMINYLTANS/FEXXIII ACETYLGUCOSAMINYLTANS/FEXXIII ACETYLGUCOSAMINYLTANS/FEXXIII ACETYLGUCOSAMINYLTANS/FEXXIII ACETYLGUCOSAMINYLTANS/F		1	П	Т	1	- I		$\top$	Homo septens DEADIH (Asp-Glu-Ala-Asp/His) box polypepted 5 (NY) I		7	٦.	
n Probes E	Top Hit Database Source	Z	TOBOSINA	MISSEL	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	SWISSPROT	SWISSPROT	EST HUMAN	NT	EST HUMAN	EST HUMAN	SWISSPROT	EST HUMAN	EST HUMAN	EST HUMAN	!	4758139 N I 30.1 EST HUMAN	SWISSPROI	SWISSPROI	
Single Exc	Top Hit Accession No.	91.2			5.0E-18 AI280214.1	44076.1	14076.1	4.0E-18 AA621814.1	4.0E-18 Q06430	4.0E-18 Q08430	1	١	3.0E-18 AL163247.4		8 Q39575	8 195406.1	19 AA281961.1	19 AA281961.1 19 AW974902.1		AW8529	19 P34986	19 P34986	
	Most Similar (Top) Hit BLAST E Value	6.0E-18 XX17		6.0E-18 P52181	5.0E-18	4.0E-18 BE0	4.0E-18 BE	4.0E-18	4.0E-18	4.0E-18					L		2 8.0E-19	9 9.0E-19	1	1.88 7.0E-19	1	١	
	Expression Signal	1 98	1	4.28	11.59	1.37	1.37	21.73	0.98	0.98	34.38	3.2	1.1	6.4			5.62	3.19	1				
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Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo sapiens Xq pseudoautosomal region; segment 1/2	DKFZp762F192 r1 762 (synchym: hmel2) Homo sapiens cDNA clone DKFZp762F192 5	Homo sepiens mRNA, chromosome 1 specific transcript KIAA0501	ENDARROMENT NIM MGC 56 Home services CDNA done IMAGE:4287674 6	DETA A ADDENICO DE PEDENCO	DE IAY AUNER CONTROL TO CONTROL T	BETA-2 ADRENEKGIC KECEPLOR	ILIM-ONLY PROTEIN 8 (TRIPLE LIM DOMAIN PROTEIN 9)	LIM-COLLY PROTEIN 6 (TRIPLE LIM DOWNIN TROTEIN 9)	AV708136 ADC Hamo agains guna quae ADCAMATTO	Homo sapiens chromosome 21 segment no 1000 l	qe91e02x1 NCI_CGAP_NIGS Homo septens curva ciune introc ie recoco commerciale polificiale del processor del pro	601304125F1 NIH_MGC_21 Homo saplens cONA cione IMAGE:3636310 9	yo79g07.7 Soares adult brain N2b4HB56Y Homo saptens cunn cione invince: 104 too o eatilise to commission	MER10 repetitive element;	Human gene for An-receptor, each 1-e	Homo saplens protein tyrosine prospriates, inchreaching type emanages of the MEDS?	849512.51 Scenes_testis_NHT Homo sapiens cDNA cione IMAGE:1383631 3 similist to contains microstrat. MER37 remotifiue element:	ISAAA ANAMOR AFARAN ANA ANAMOR Homo saplens cDNA	TWITH SUBSAMIN 1 SEOLIENCE CONTAMINATION WARNING ENTRY	ACC 3015 MILL MGC 72 Home seplens CDNA clone IMAGE:3916231 5	AV725422 HTC Homo serviens cDNA clone HTCBTA01 6	Homo saniens chromosome 21 segment HS21C047	Human RXP21 mena	TO EARTORY DECEDITOR IKE PROTEIN 14	Action of Scores present thems NoHPU Home septens CDNA clone INAGE:484895 3" similar to	2030012.51 Source program. Contains L1.03 L1 repetitive element;	Г	P97461 40S RIBOSOMAL PROTEIN SO.; SONA clone IMAGE:940087 similar to TR:G1224088		Т	
	Top Hit Database Source	K	EST HIMAN		1400 11 11 11 11 11 11 11 11 11 11 11 11 1	ESI HUMAN	SWISSPRO	SWISSPROT	SWISSPROT	SWISSPROT	EST HUMAN	N	EST_HUMAN	EST HUMAN	·	EST HUMAN	NT	N.	NAME OF THE PARTY	NOWOLL TOT	ESI HUMAN	SWISSPRO	EST HOMAN	ESI DOMAN	1	- N	SWISSPROI	EST HUMAN		EST_HUMAN	EST HIMAN		EST HUMAN
STOLL TOWN BIRLING	op Hit Acession No.	JZ71735.1		T		27.1		228997				AL163201.2	Al311783.1			H30795.1	D38044.1	4758977 NT			5.1	P39188	BE622434.1	AV725123.1	AL163247.2	U03888.1	P23273	A A N 3 7 8 1 8 1		AW303868.1	A A 64 65 5 6 4	AA010990.1	AA616335.1
-	Most Similar (Top) Hit BLAST E Value	8.0E-19 A.		- L	•	_		3.0E-19 Q			3.0E-19 A	2.0E-19 A	2.0E-19			1.0E-19	1.0E-19			1.0E-197	7.0E-20 E				4.0E-20/	3.0E-20	3.0E-20	9 OE 30	_	2.0E-20	L	2.0E-20	2.0E-20
-	Expression Signet	1.38		<u>\$</u>	re.0	+	0.98	96.0	1.07	1.07	1.25	24.97	1.37	1.58		1.18	201	545			0.74					1.22	1.48	4.07		48.48		3.05	3.05
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	Econ SEQ ID NO:	0790	3	9845	6584	7573	8765	8765	9158	9158	9316	7457	0345	5515	3	7081		1	1_	8327	8217	8486	9145	9454	6578	7060		L	19481	5840		6101	6101
	Probe SEQ ID NO:	1000	8	4865	999	2611	3762	3762	4163	4163	4324	2489	66,	37.4	2	200	2845		2//4	3317	3202	3478	4150	4464	1581	2079	4088		\$ 26 26	820		1094	1094

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	724610.71 NCI_CGAP_Ut/4 Homo septens cDNA clone IMAGE:2761086 3' similar to SW:1436_MUUSE P97461 40S RIBOSOMAL PROTEIN S5.;	ZONADHESIN PRECURSOR	ZONADHESIN PRECURSOR	Homo sapiens matate denyungarase 1, Ivan (souther) (men-	A11406.11 NCI_CGAP_GCB1 Home explans guith circle into can be considered to the constant of th	hre4b06.x1 NCI_CGAP_Kid11 Homo sapiens cDNA done IMAGE:3135106 3 simular to contains Link Link Link Link Link Link Link Link	1 a horaz 4.4 Selected chromosome 21 cDNA library Home sectors cDNA clone MP1p12-8/21	COUNTY OF THE STATE OF THE STAT	LAMININ DELTA COLAND DESCRIPTION OF AMININ CHAIN BS)	LAMININ BETA-2 CHAIN TRECONOCA (9-CAMININ) COMMINING COM	266/806.T Soares pregnant uratus replace Anna done IMAGE-3838310 5	601304125F1 NIH MGC_ZI INGIN SEQUENCE CONTROL (PTPN21) MRNA	Home sapiens protein growing principal and a close IMAGE:3933880 5	601649871F1 NIH MGC /4 name square control meson in the 21 (PTPN21), mRNA	Homo sapiens protein principal contract from 1971	Homo sapiens melarioma arrugari, raminy C., 1 (1970-1573) April 1970-1973 April 1970-1970 Apri	PMS3 MRNA; contains OFR.tt OFR repetitive element;	2415406.s1 Stratagene fetal retina 837,202 notice satisfies CONN constitutions	Homo septens chromosome zi sogment in zince	Home saplens LGMD/25 gene Home sariens cDNA	UNS-HIU40001/VZUU-0509 IZ III Opposite III O	Home salars man for KIA40397 protein, partial cds	Figure   F	TANADHERIN PRECIRSOR	JONADHESIN PRECIESOR	HAMAGAR #1 NCI CGAP Pr4 Homo septens cDNA done INAGE:1043718 similar to contains MER29.b2	EST_HUMAN   MER29 repetitive element;	ar88d12.x1 Barstsad colon HPLRB7 Homo saprens curvo curo invocato de signifar to TR-015408 Q15408		1
Top Hit Database Source	EST_HUMAN	SWISSPROT	SWISSPROT	٦	EST_HUMAN	_	ESI HUMAN	ES HOMAN	SWISSPROT	SWISSPROT	EST_HUMAN	EST HUMAN	Ę	EST_HUMAN	FZ.	LZ	EST HUMAN	EST_HUMAN	Į,	F	EST HUMAN	Į.	I Z	EST HUMAN	SWISSPRO	SWISSPRO	EST_HUMAN	EST HUMAN.	EST HIMAN	
Top Hit Acession No.	2.0E-20 AW303868.1			5174538 NT	1.0E-20 AA281961.1		1	<u></u>		1		3E408611.1 .	6902031 NT	3E968839.1	5902031 NT	4885474 NT	AAB70713.1	AA218891.1	AL163201.2	AJ007973.1	BE163247.1	AB007857.2	AB007857.2	BE064410.1	028983	Q28983	AA567657.1	AI801264.1		M102450.1
Most Similar (Top) Hit BLAST E Value	2.0E-20.A	2.0E-20 Q28983	2.0E-20 Q28983	2.0E-20	1.0E-20.A		1.0E-20 B	9.0E-21 /	7.0E-21 F	7.0E-21 F	7.0E-21	6.0E-21	6.0E-21	5.0E-21	5.0E-21	5.0E-21	4.0E-21	3.0E-21	3.0E-21							2.0E-21	1 0F-21			9.0E-22
Expression Signal	36.55	4.58	4.58	1.73	2.2		1.12	1.16	1.61	1.61	6.12	0.98	67.0	3.34	0.83	6.35	1.66	1.05	0.99		18.78			2.38		2.35	14	9		1:1
ORF SEQ ID NO:	<b>†</b>	14756	14757		1.20K4		14293		12102			13971	L		10964		11787	L	L			10969	10970	60	12648	12649	44070	1		8 14266
Exan SEQ ID NO:	OF 82	2773	9773	9948	1	3	8308	7862	6699	6889	9128	8985	5929	9232	5929	9642	<u> </u>		L	L		1		L	L			0239 6275	1_	8278
Probe SEQ ID S	33.45	4780	4789	4969	Ş	3	4316	2842	2016	2018	4133	3987	913	4738	4548	4657	260	1801	2562	3008	145	922	922	1195	2587	2567		1236	2/2	4286

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Care Litrate 281088-078-h05 HT0179 Home septems cDNA	United services chromosome 21 segment HS21C046	TOTING SEAMEN STATE OF THE CURSOR (ALPHAZM)	ALPHA-Z-WACKOULOCKIN TO COMPIE COMPIETE COS	Holin Sayana Sanardan Instituti 182	Homo Sapiteris Al posture and the service ANA Acros IMAGE 2158611 S' similar to gb:L19593 HIGH	tm4410x1 NCI_CGAP_CO1+ runts aspects and AFFINITY IL1 IL1 repetitive element;  AFFINITY INTERLEUKIN-8 RECEPTOR B (HUMAN);contains L1.11 L1 repetitive element;  AFFINITY CARD BUSE Homo seniens cDNA clone IMAGE:2429839 3' siniliar to SW:RL21_HUMAN	WIGGOLA XI NCI_COSMDILIZOREN L21.; P46778 608 RIBOSOM LIVIGA Palated mana	Human diromoscular process and the second se	optable XII States Inspective element; contains MER12.t2 MER12 repetitive element;	W/3405.81 Soares meaningle Zyta in resident	Invited in the Charles AMP-activated gamma 3 non-catalytic subunit (PRKAG3), mRNA	Homo septens procent misses; viiii DMA-ST0262-281189-001-412 ST0262 Homo septens cDNA	PARA SN0020-010400-009-h02 SN0020 Homo saplens cDNA	Human familial Abheimer's disease (STM2) gene, complete cds	Human DNA, SINE repetitive element	AVB47248 GLC Homo saplens cDNA clone GLCAWC07 3'	Ratus novegicus RIM1B (Rim1B) mRNA, complete cds	Homo septens chromosome 21 segment HS21C049	Rettus norvegicus RiM1B (Rim1B) mRNA, complete cds	Homo saplens KIAA0851 gene (partial), XT3 gene and LZ1PL1 gene	Human matrix Gla protein (MGP) gene, complete cas	TENASCIN-X PRECURSOR (TN-X) (HEXABRACHION-LIKE)	TENASCIN-X PRECURSOR (TN-X) (HEXABRACHION-LIKE)	19373111.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:1943/6/ 5 SILILIE TO 19573/6/ 5 SILILIE TO 1957311.X1 NCI_CGAP_PR28 HOMO score CDNA clone		Т	Π.	Т	Homo saplens chromosome 21 segment HS21C010	
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-	増せる	Væue	8.0E-22	7.0E-22	7.0E-22	7.0E-22	4.0E-22	3.0E-22 A	3.0E-22	3.0E-22	3.0E-22	2.0E-22	2.0E-22	2.0E-22	2.0E-22	1.0E-22	1.0E-22	1.0E-22								7 2.0E-23	7 2.0E-23				1.0E-23	_
	Expression Signal		5.16	4.6	211	0.98	13.85	0.88	2.28	125	3.17	209	44.	3.8	1.53	1.41	•	1.7		1.59				3.75		1.37	1.07	4.12		5 2.39		4.78
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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor  Top Hit Descriptor  Top Hit Descriptor	88/18/20 ST STRING ST PROTEIN ; TRE-18922 E19822 CA PROTEIN ; OLFACTORY RECEPTOR-LIKE PROTEIN IS	OLFACTORY RECEPTOR-LIKE PROTEIN IS	QV0-DT0047-170200-122-e08 D1004/ Hanto septems CDNA clans DKFZp434A2311 5/	DKFZp434A2311_71 434 (synonym: nieso) India openieso cds	Macaca fusceta mRNA for Testis-Specific Troops (2015)	Home saplens chromosome 21 segment 102 100 100 Homesome 21422, segment 3/3	Homo septens sos in contra contra contra Homo septens contra clone o-zrc08	HSCZRCX61 normalized filters to the company of the	401109.11 Strangelin in the Strangeline CONA	RC3-NN0068-05000-021-05-05-05-05-05-05-05-05-05-05-05-05-05-	Homo saplens CGI-127 protein (LOCs10+0), many homo saplens cDNA	QV0-ST0284-100400-185-610 ST0284 Franks Cd8	Mus musculus mRNA for HG I Kereun, per var co	Homo saplens PTEN (PIEN) gene, and a form IMAGE:911764 similar to contains MER1.02	ne92e10.s1 NCI_CGAP_Kld1 Homo sapiette Control ne92e10.s1 NCI_CGAP_Kld1 Homo sapiette	MER1 repetitive dement : II.3-CT0219-161199-031-D04 CT0219 Hono saptens cDNA clone IMAGE:121783 5	Т	1			Homo saplers hypometical process. Homo saplers in the same saplers in the same saplers in the same saplers in the same saplers in the same saplers in the sa	٦	- 1	T	Т	Т	Т	٦	- 1	1	Talio odport	
Flores	Top Hit Database Source	EST HUMAN	Ţ	Т	T	1			EST HUMAN	EST HUMAN	FST HUMAN		EST HUMAN	LN	Į.		EST HUMAN	EST HUMAN	FST HUMAN	EST HUMAN	N.	F	NT	EST HUMAN	SWISSPROT	SWISSPROI	SWISSPROT	EST_HUMAN	ĮN.	SWISSPROT	EST HUMAN	Z	
Single Exon Propes Express	HR Acession No.		T	7,79	T	T				-	T	2.0E-24 AW 896109.1				T	$\neg$	AW850271.1	198107.1	RE170857 1	23321	TN 1253321 NT		BE88801	P17008	P17008		AL04022	9635487 NT	206055	5 BE162737.1	6 AL 163218.2	
-	Most Similar (Top) Hit BLAST E Value	9.0E-24 AA663213.1	8.0E-24 PZ	8.0E-24 PZ3209	7.0E-24 AWB3/804.1	7.0E-24 ALUSB459.1	6.0E-24 AB001421.1	E OE 24 A 1229043.1	9.0E-24 F08337.1	A AC 30 C	Z.VE-2-10	2.0E-24 A	1.05-24	1.05-24	1.0E-24 D00423.	1.0E-24	7.0E-25	8.0E-25	4.0E-25	4.0E-20	4.0L-25	3.0E-25	2.0E-25	2 OF 26								9.0E-26	
	Mos Expression (T	. 2.52	1.13	1.13	0.83	1.18	23	33.01	8.31	20.1	1.8	0.78	2.5	1.1	4.21	1.8	3.17	12	1.54	2.89	3.14	3.16	3.10	3.48	6.00 86.00	4 60	3.4	25.0	200				
		+-	282	14483		H	H	10885	13869	14871	12379	H	11725	-	12988	-	14798	11672	11478		-	13280	13281	11375	1234	2007	100	14040	10423		12458	14679	12302
	ORF SEQ ID NO:		14482												L					28			8259		7222	\$ <del>5</del>	888	0900	5412	6228	134	5695	7383
	Eson SEQ ID NO:	CRAR	503	9503	8784	8968	5720	5848	8865	8897	7261	L	1_	L						Ŀ	9183		L		۱ ۱			4066 90	363		2367 7		2412 7
4	Probe SEQ ID S	3	8 8	4543	2 18	289	8	828	3863	4919	2285	3709	1657	2802	2054	4448		4637	1424	3318	4190	3246	3248	1329	2245	2758	4066	\$		12	Ä	4	7

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su87h08.x1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2783295 3' similar to gb:K00559 au87h08.x1 Schneider fetal brain 00004 Home saplens cDNA clone IMAGE:2783295 3' similar to gb:Ko0558 2030/10.r1 Stratagene colon (#937204) Homo sapiens cDNA clone IMAGE:588427 5' similar to TR:G695374 zo30f10.r1 Strategene colon (#937204) Homo sepiens cDNA clone IMAGE:588427 6' similar to TR:G695374 Homo seplens glyceraldetryde-3-phosphate detrydrogenase (GADPH) mRNA, complete cds w/49c04.x1 NCI\_CGAP\_Lu19 Homo seplens cDNA clone IMAGE:2406160 3' similar to contains THR.b2 2730d08.11 Stratagene neuroepithelium NT2RAMI 937234 Homo sepiens cDNA clone IMAGE:548943 6 Homo sapiens chromosome 9 duplication of the T cell receptor beta locus and trypsinogen gene families 2352/104.1 Stratagene neuroepithsilum (#837231) Homo sapiens cDNA clone IMAGE:845271 6. as38h08.x1 Barstead sorta HPLRB6 Homo sapiens cDNA clone IMAGE:2319519 3' similar to TUBULIN ALPHA-1 CHAIN (HUMAN);

PM2-SN0018-220300-002-407 SN0018 Homo sepiens CDNA
ADP,ATP CARRIER PROTEIN, LIVER ISOFORM T2 (ADPIATP TRANSLOCASE 3) (ADENINE
NUCLEOTIDE TRANSLOCATOR 3) (ANT 3) Homo septens X-linked anhidroffic ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat as38h08.x1 Barsteed earta HPLRB6 Homo septens cDNA clone IMAGE:2319519 3' stmilar to DKFZp566L171\_s1 566 (synonym: hfkd2) Homo sapiens cDNA clone DKFZp566L171 3' similar to gb:M14338 VITAMIN K-DEPENDENT PROTEIN S PRECURSOR (HUMAN); H. sepiens DNA for endogenous retrovirsi tike element hd02e12x1 Soares\_NR\_T\_GBC\_S1 Homo sepiens cDNA clone IMAGE:2908366 3\* MR2-BN0114-240500-030-607 BN0114 Homo septens cDNA QV4-HT0538-020300-123-e02 HT0538 Homo saplens cDNA **Top Hit Descriptor** M.musculus mRNA for astrocytic phosphoprotein, PEA-15 Homo saplens chromosome 21 segment HS21C027 G695374 THYROID RECEPTOR INTERACTOR: Home septens chromosome 21 segment HS21C082 G695374 THYROID RECEPTOR INTERACTOR WP:F49C12.11 CE03371; Human DNA, SINE repetitive element Single Exon Probes Expressed in HBL100 Cells THR repetitive element EST HUMAN SWISSPROT EST\_HUMAN EST HUMAN EST HUMAN EST HUMAN EST\_HUMAN EST HUMAN EST\_HUMAN EST HUMAN HUMAN EST\_HUMAN EST\_HUMAN HUMAN Top Hit Detebase Source EST EST E 둗 눋 8.0E-27 AW162737.1 8.0E-27 AW864776.1 8.0E-27 AW162737.1 Top Hit Acesslon 1.0E-26 BE814995.1 1.0E-26 AF261085.1 8.0E-27 AI831462.1 3.0E-26 AA152464.1 AL163282.2 AL038099.2 5.0E-26 AI708235.1 3.0E-26 D14547.1 3.0E-26 AA115895.1 3.0E-26 AA152464.1 8.0E-27 P12236 7.0E-26 X89211.1 7.0E-26 AW340153.1 6.0E-26 AF028308.1 5.0E-26 AI708235.1 AF003528.1 AA206131. 8.0E-27 2.0E-26 1.0E-26 2.0E-26/ 2.0E-26 8.0E-26/ 7.0E-28 (Top) Hit BLAST E 268 83.09 83.88 273 7 1.04 57.21 7 5.99 234 5.27 1.25 278 1.78 6.17 5.17 1.32 1.78 Expression 13147 12199 11441 <u>4</u> 10075 13199 10218 12575 13698 13697 11196 11197 12261 13879 ORF SEQ ID NO: 8127 8386 7083 6388 5091 5587 5698 8695 5203 8695 6821 767 6716 88 6162 7143 8283 6162 6540 8874 9031 SEQ ID ÿ 3111 243 1391 137 2615 553 1391 11 55 153 153 3160 3691 1158 3691 1158 <del>1</del>981 1721 3873 2164 4035 327 Probe SEQ ID 혖

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Human endogenous retroviral element HC2	HIGHT2-X1 Soares_NFL_T_GBC_S1 Homo eapliens CDNA clone IMAGE_297.001	078040 ORF2: FUNCTION UNIXNOWN.:	R. ratus KTA3 minuto for province of the sapiens con A	PM0-B105Z/-090100-01-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1	Home sapiens alpha NAC mixture, company chara IMAGE:1000699 similar to gb:M17886 60S	nkotbto.st NCI_CGAP_Pri1 Homo saprens Color Constant Color C	ACIDIC RIBUSCIMAL TAND LENGTH HOME SAPIENS CONA Clone IMAGE 2976879 S' similar to TR: 076940 histhritz. I Seares_NFL_T_GBC_SI Home sapiens CONA clone IMAGE 2976879 S' similar to TR: 076940	076040 ORF2: FUNCTION UNKNOWN: .	Homo sapiens Jun dimerization protein gene, parter des, consistent des and unignown gene	Homo septens Jun dimerization protein gene, partial cds, dos gene, compressional desperanta of the companies chromosome 21 segment HS21CO46	Homo sapiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (ULEC1, UNO LD, CICCIE)	complete cds) Intogeo1.x1 NCI_CGAP_Kid13 Homo septens cDNA clone IMAGE:3146256 3' similar to contains MER29.b3	MER29 repetitive etement;	SECRETED NEUREXIN III-ALTHA-C PRECONOR 1727 173 174 174 174 174 174 174 174 174 174 174	AU142750 Y79AA1 Homo saplens clura 1/2000 INAGE:2455652 3' similar to contains THR.b1	wo18007.xt NC_CGAP_ram name servers	1HK repairing scanned in No2-IP Homo septems cDNA clone IMAGE:148443 b  V89f10.r1 Soeres placents Nb2-IP Homo septems cDNA clone IMAGE:2895504 3' similar to SW:GG95_HUMAN				Г	Т	1		
	Top Hit Databese Source			EST HUMAN	П	EST HUMAN			EST HUMAN	EST_HUMAN					EST HUMAN	EST HUMAN	EST HUMAN		EST HUMAN		EST HUMAN	<b>EST HUMAN</b>	_	NT EST HIMAN	101	Z	
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ŀ		-	7.0E-27/270	7.0E-27 AV	15 15	16		<u>:</u>	20E-27 A	A IZ	2.0E-27 A	E-27/A	E-27	1.0E-27	1.0E-27	9.0E-28	9.0E-28	/,UE-20	5.0E-28	3	4.0E-28	4.0E-20	2	3.0E-28	20E-2	2.0E-2	
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			8/2		888	6975	9144	5122	448	88	2	3	6474	500g	98	5206	5364	6166	5371	8902	7517	7927	8052	۽ ا	7070	919	200
		<u> </u>	676	_		1880	4149		1855	1	27.6	3 9	318	} · 8	g 99	3	100	1162	316	3802	26.53	2008	3035	1	128	6	147
	Probe SEQ ID	Ö	100		4	18	4				<u>L</u>		$\mathbb{I}$				_	_	<u> </u>	_		L	1		_1		لـ

														4"			*	<u> </u>	=	+		Ť	í i	-	٣	7~	F		
Page 85 of 209 Table 4 Single Exon Probes Expressed in HBL100 Cells	Top Hit Descriptor  Top Hit Descriptor  Top Hit Descriptor  Top Hit Descriptor	repetitive element : Hamo septens chromosome 21 segment HS21 C009	Human gene for Ant-received, and the supers con Anterior con Anterior Anterior and the supers con Anterior Anterior and the supers con Anterior Ant	QV1-B 10621-125000 Limen zinc finger probein ZNF131 mRNA, perdel cds	O FACTORY RECEPTOR 15 (OR3)	EST378521 MAGE resequences, MAGI Homo sakkeis Communication in the INAGE 3356387 6	601114990F1 NIH MGC 10 Hours septens cONA clone IMAGE:2466985 3' similar to INC 1970	wp69b01x1 NCI_COXT_DILET   Contains LTR7.b1 LTR7 repeating desired:	O15475 UNIVAMED 11-12 Homo sepiens chromosome 21 segment HS210003	t-	1	Homo saplens envelope protein Kilcho (env) gene, complete cds	Homo saplens envelope protein INCO (2017) and Homo saplens CDNA clone IMAGE: 2492663 3° smilistra in INCO (2017)	wre5d10x1 NCL_CCAL_CTAL TR. O16546 O16548	7		+	7	٦	7	1_	Т		П	Т		TI		
Page 8 Tal	Top Hit Database Source	EST_HUMAN		HUMAN	Todage	SWISSPROI	EST HUMAN	_	EST HUMAN		EST HUMAN		Į.	-	EST HUMAN		NT TOWN	EST HUMAN	IN	Į.	EST HUMAN	ESI HOMON	EST HUMAN	EST_HUMAN	EST HUMAN	TOT LIMAN	NT IN	ļ ——	-
xou	- a "	EST	토	183	킥	80		+		4	<u>S</u>	T	T	T	쀠			١						-	٠-		_ -		
Single E	Hit Acession No.	48634.1	20E-28 AL 103208-2	1.0E-28 BF333236.1	9410.1	1.0E-28 PZ3275	7.0E-29 AW966447.1	7.0E-28 DE CONTON	6.0E-29 AI936748.1	1163203.2	4.0E-29 AI752367.1	B042297.1	2.0E-29 AF084869.1	1-004003.	2.0E-29 AI963604.1		2.0E-29 AI963604.1	BE091133.1	X51755.1	D25303.1		BE008028.1			AWB37471.1		10 AI338551.1		
	-7	2.0E-28 A1348	20E-28 AL 103 cus.	2 8 PR	28	-28 P2	8 8	-	-28 N	6.0E-29 AL1	E-29 A	E-29 A	E-29		DE-29		SE-20	2.0E-20.2	8 OF 30	6.0E-30	6.0E-30	3.0E-30		5.0E.30	10 V	3	3.0E-30		
	Most Similar (Top) Hit BLAST E Value	20E			18	1.8	9		90.0	6.0	4.0	မ္မ	20	۲	ัล		7				1_	1			1		<u> </u>		
.		1.68	0.68	<u> </u>	222	1.53	12.	1.34	99.6	1:37	1.75	22	1.15	1.15	4 68		4.68	2.09	4.34	200	545	18		40.12	3	=	3.35	0.82	
	Expression Signal				١										<u>_</u>		- <del>-</del>	4	$\dashv$	4		2	22	2	72	2		13680	
		12496	13318	11504	226	9707	11627		0830	3		1,1970	10532	10533		188	11554	14134			11808	13152	ह्	13904	12172	12173		2	
·	ORF SEQ ID NO:	\	L	Ц				L			-		8282	2322 E525	1-	6499	6400	9152	6481	6532	6731	8133	8133	8906	7063	7083	8139	1188	
	SEQ ID	22.5	2 88	6445	7137	9426	8974	١	١	5621 9823				$\perp$				L		L			Ш	3006	L	1 78 K	7000	3672	
	Probe S		2000	448	2158	4436	5003	1001	34/0	88	200	3161	4280	\$ 1	489	1501		1901	484	150	1=	     	4		Ϊ	1	L	ľ	

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	CA/0-CT0307-310100-158-h03 CT0307 Homo septens cDNA	HSC23F051 normalized infant brain cDNA Homo septens cDNA done c-23106	RC5-HT0582-110400-013-H08 HT0582 Homo saplens cDNA	IL2-NT0101-280700-116-E04 NT0101 Hamo sepiens cDNA	Homo saplens Y-linked zinc finger protein (ZFY) gene, complete cds	UI-H-BI1-#fo-c-12-0-UI.s1 NCI_CGAP_Sub3 Hamo sapiens cDNA clane IMAGE:2722558 3'	601119860F1 NIH_MGC_17 Hamo saplens cONA clane IMAGE:3029438 5	601119860F1 NIH_MGC_17 Homo septens CDNA clone IMAGE:3029438 6	C18939 Human placenta cDNA (TFujiwara) Homo saplens cDNA clone GEN-570C01 6	hd30b04.x1 Soares_NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE:2910991 3' similer to contains MER1.t3 MER1 MER1 repetitive element;	Homo sepiens chromosome 21 segment HS210003	ac77b08.s1 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:868599 3'	602022560F1 NCI_CGAP_Bm67 Hamo saplens cDNA clone IMAGE:4157891 5	EST186868 HCC cell line (matastasta to liver in mouse) if Homo saplens cDNA 5' end	Homo sapiens hypothetical protein FLJ20420 (FLJ20420), mRNA	Hamo seplens chramosome 21 segment HS21C008	OLFACTORY RECEPTOR 16 (OR3)	OLFACTORY RECEPTOR 16 (OR3)	EST84555 Colon adenocarcínoma IV Homo aptiens CDNA 6 end	hw05a11.x1 NCI_CGAP_Lu24 Homo seplens cDNA clone IMAGE:3182012.31	hw05a11 x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3182012 3	Homo sepiens calcium channel apha1E subunit (CACNA1E) gene, exons 7-49, and partial cots, etiomatively spliced			-1/2	Homo sapiens chromosome 21 segment HS21C080	Homo saplens SET domain and mariner transposase fusion gene (SETMAR) mRNA	Homo sapiens SEC83, endoplasmic reficulum translocon component (S. cerevisiae) like (SEC83L), mRNA		П	DKFZp761G1513_f1 /01 (synonym: namyz) monto equate consistent consistent consistent of the consistent
Top Hit Database Source	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	_ 1	EST_HUMAN	EST_HUMAN	EST_HUMAN	<b>EST_HUMAN</b>	EST_HUMAN	_	EST_HUMAN	EST_HUMAN	EST_HUMAN	T	NT	SWISSPROT	SWISSPROT	EST_HUMAN	EST_HUMAN	EST_HUMAN		NT	Z L	F	NT.	<u> </u>	 =	<b>EST HUMAN</b>	EST HUMAN	EST HUMAN
Top Hit Acession No.	16.1	-	7.1		2.0E-30 AF114156.1 NT		Γ		1	x97.1	1L163203.2 NT	77.1		45.1	8923389 NT	38.2			7.0E-31 AA372637.1 E		7.0E-31 BE326517.1 E	8.0E-31 AF223391.1	Γ	2	+	80.2	5730038 NT	F005871 NT	1	88.1	245.1
Most Similar (Top) Hit BLAST E Vatue	2.0E-30 AW8573	2.0E-30 F08688.	2.0E-30 BE1768	2.0E-30 BE7652	2.0E-30/	2.0E-30/	2.0E-30 E	2.0E-30	1.0E-30 C18939.	1.0E-30 AW468	1.0E-30 AL1632	1.0E-30/	1.0E-30 BF3477	1.0E-30	8.0E-31	8.0E-31	8.0E-31 P23275	8,0E-31 P23275	7.0E-31	7.0E-31	7.0E-31	6.0E-31	5.0E-31 M60694	5.0E-31	4.0E-31 AJ2717	4.0E-31 AL1632	4.0E-31	3.0E-31	2.0E-31 AW838		2.0E-31 AL119;
Expression Signal	121	2.48	8.55	6.5	6.88	223	1.61	1.61	13.84	3.42	3.15	2.78	1.95	0.79	6.71	23.47	1.01	1.01	2.43	2.15	2.15	2.5	3.17	3.17	3.18	1.4	1.63	1.54	2.35	1.01	1.52
ORF SEQ ID NO:	10701		11505	12716	12885	13707			10355	10571	10746	12248	12486		11100		14733	14734		12676	12676		10268		L			12604			12373
SEQ ID	6692	878	8448	7603	7869	8704	9618	9618	5342	5568	5730	7128	7384	7998	6908	7321	L	9747	L	L		P.OBG	L	L			Ľ		L	L	Ш
Probe SEQ ID NO:	198	1088	1449	2843	2846	3700	4633	4633	284	683	706	2150	2383	2882	1080	2347	4763	4783	é	2597	2597	960	3 6	ģ	ğ	47B2	2716	2610	1873	2162	2278

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	asB8f11.s1 Stratagene fetal retins 937202 Home suprens Colors Colors Colors Colors Colors Stratagene fetal retins 937202 Home suprens Colors Stratagene MAGE-B2 (MAGE-B2), MAGE-B3), MAGE-B4 (MAGE-B4), and MAGE-B1	Moderally genes, complete ods Or EACTORY RECEPTOR 2C1	OLEACTORY RECEPTOR 2C1	OLFACTORY RECEPTOR 2C1	DKFZp547B235_r1 547 (synonym: htbr1) Home saplens cONA come DN report B223 5	DKFZp547B235_r1 547 (synonym: hfbr1) Home eaplens cDNA clone DNA clone IMAGE-16753843'	oz15a09.x1 Soares fetal liver spleen_INFLS_S1 Homo saprens Curva cluire inva-	Homo saplens PRO1181 mRNA, complete cds	Homo saplens chromosome 21 segment HSZ1CV4o	Homo sapiens FLH gene, partie	AV731500 HTF Homo sapiens cDNA clore HI I FANCAL O	Homo sapiens mRNA for phenyalanyi RNA synthetiase, conjugate we	601573207F1 NIH_MGC_9 Hano sapiens cDNA clane IMAGE:3001453 5. 1miles in TR-088539 088539	INVOTCOSAT NCI_CGAP_Lu24 Hamo sapiens cDNA clane IMAGE:3162210 3 Blilling 20 11	WW DOMAIN BINDING PROTEIN 11.:	Hours september chain alcohol dehydrogenase family member (HEP27) mRNA	Home septents structure to contains OFR.ft OFR Than 5 septents OFR.ft OFR Than 5 septents OFR.ft OFR Than 5 septents OFR.ft OFR Than 5 septents OFR.ft OFR Than 5 septents OFR.ft OFR Than 5 septents OFR.ft OFR Than 5 septents OFR.ft OFR Than 5 septents OFR.ft OFR Than 5 septents OFR Tha	repetitive element;	AV /30036 R. I' TIMIN September CON CONTRACTOR 5	AV 300 10 HT FIRST INSECTIONS MAGE Home sapiens CDNA	Lorno seriens chronosome 21 segment HS21C085	OV1-FT0169-100700-271-e02 FT0169 Homo sepiens cDNA	Homo saplens solute carrier family 5 (choline transporter), member 1 (2000),	Homo sapiens spermidine synthase (SRM) mRNA	Homo sapiens spermidine synthase (OKM) Illinoids	Homo sapiens chromosome 21 segment nozi coco	Homo sapiens mRNA for KIAA0599 protein, parted to CA segments), 5' end	Human TCR variable region Vs30 Subramily gene (VAV) C.1	Homo saplens chromosome 21 segment HSZ1CoV7	
	Top Hit Detabase Source	EST_HUMAN	NT	SWISSPROI	SWISSPROI	ENT HIMAN	EST HIMAN	EST HUMAN	Į.	Ę	LZ	EST HUMAN	IN	EST HIMAN		EST HUMAN	N	Ę	EST HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	EST HIMAN	NT	L N	BNT	Ż	Z	L <sub>N</sub>	Z	
	p Hit Acession No.	158824.1		1		1	1	T	148877 4	T	7202 4	724500 4	3.0E-32 AV/31500.1	1.0E-32 D84430.1	E/4328.1	32711	5031736 NT	5031736 NT	AI590115.1	7.0E-33 AV730056.1	7.0E-33 AV730015.1	7.0E-33 AW971307.1	AL163285.2	BF3/3010.1			A1 48228			AL 163207.2	
-	Most Similar (Top) Hit BLAST E	2.0E-31 AA	1.0E-31 U93163.1	1.0E-31 095371	1.0E-31 095371	1.0E-31 095371	1.0E-31 AL1343/0.1	4.0E-31 AL 1343/0.1	0.0E-32	0.0E-3Z	4.0E-32	3.0E-32 11	3.05-32	1.0E-32.L	1.0E-32	9.0E-33 Bi	7.0E-33	7.0E-33	7.0E-33 A	7.0E-33	7.0E-33	7.0E-33	6.0E-33 A	5.0E-33	5.0E-33	0.05-33	3.UE-33	0.00-20	0.0E-30	5.0E-33	1
	Expression (	4.03	10.14	9.05	9.05	9.05	1.26	1.26	5.49	84.69	1.7	2.46	57.22	1.49	1.65	522	11.74	11.74	1 03	7.4	101	14.85	0.8	1.43	0.97						1.0
	ORF SEQ ES	12465	10079	11686	11687	11688	14467	14468	12113	11057		10505	11483				40144			12184	44707		Flee	2			7 11935			15001	[]
	SEO D NO:	7346	2002	8620	6620					6027	5934	5490	6424	7566	8037	97,0	١	5142			200				<u> </u>	58 6847	58 6847	10 7187	3945 8943		1111 6117
	Probe SEQ ID NO:	8,8%	2 4	1823	1623	1623	4501	4501	2024	1017	918	453	1427	2604	3020		3	62		2097	25/6	/6/2	3852	1740	1842	1858	1858	2210	18	8	=

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Homo septens RAB1, member RAS oncogene family (RAB1) mRNA	àb51b11.r1 Stratagene lung cardinoma 897218 Horino saplens cDNA ctone IMAGE:844317 5° similar to contains Alu repetitive element:oontains MER28.b2 MER29 resettitive element:	Homo sapiens chromosome 21 segment HS21C010	UI-H-BI2-ahl-c-03-0-UI.st NCI_CGAP_Sub4 Homo sepiens cDNA clone IMAGE:2727149 3'	ht09g01.x1 NCI_CGAP_Kid13 Homo sapiens cDNA clone IMAGE:3146266 3' similar to contains MER29.b3 MER29 repetitive element;	ht09g01.x1 NCI_CGAP_Kid13 Homo saplens cDNA clone IMAGE:3146256 3' similar to contains MER29.b3 MER29 repetitive element:	AV647851 GLC Homo septens cDNA clane GLCBCF09 3'	qb87g03.xf Soeres, fetal_heart_NbH119W Home sapiens cDNA clone IMAGE:1705204.3' similar to contains OFR.tf OFR repetitive element;	qb67g03.x1 Soeres, fetal_heart_NbHH19W Home sepiens cDNA clone IMAGE:1705204.3' similar to contains OFR.t1 OFR repetitive element;	MR0-HT0405-160300-202-d08 HT0405 Hano sepiens cDNA	ab51g11.r1 Stratagene lung carcinoma 837218 Homo sapiens cDNA cione IMAGE:844388 6' similar to gb:X00734_cds1 TUBULIN BETA-6 CHAIN (HUMAN);	Homo sapiens hypothetical protein SIRP-b2 (SIRP-b2), mRNA	Homo sapiens hypothetical protein SIRP-b2 (SIRP-b2), mRNA	Homo sapiens X-linked anhidrolite ectodermal dysplasia protein gene (EDA), excn 2 and flanking repeat regions	QV2-BT0258-071289-019-g07 BT0258 Hamo seplens cDNA	yd15e05.r1 Sceres fetal liver spleen 1NFLS Homo sapiens cDNA cione IMAGE;108320 5	Human G2 protein mRNA, partial cds	Human G2 protein mRNA, partial cds	Homo sapiens Npw38-binding protein NpwBP (LOC\$1729), mRNA	Human splicing factor SRp55-1 (SRp-55) mRNA, complete cds	tt94c08,x1 NCI_CGAP_Pr28 Hamo sapiens cDNA clone IMAGE:2249184 3*	Homo saplens hypothetical protein FLJ10989 (FLJ10889), mRNA	Homo septens splicing factor 3a, subunit 3, 60kD (SF3A3), mRNA	ADP, ATP CARRIER PROTEIN, LIVER ISOFORM T2 (ADP/ATP TRANSLOCASE 3) (ADENINE NUCLEOTIDE TRANSLOCATOR 3)	Homo sapiens X-linked anhidroitic ectodermal dysplasta protein gene (EDA), exon 2 and flanking repest regions
Top Hit Database Source	٥	EST HUMAN	Z	EST_HUMAN	EST HUMAN	ST HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	5	11	IN	EST HUMAN	EST_HUMAN	된	NT	누	NT	EST_HUMAN	11	17	SWISSPROT	IN
Top Hit Acession No.	4758987 NT	AA828821.1			E350127.1	E360127.1		1160189.1	1160189.1	E159039.1	A626683.1	11421332 NT	11421332 NT	F003628.1	BE062570.1	T70845.1		U10991.1	7706500	U30883.1	AI804667.1	8922807 NT	5803166 NT	27236	AF003528.1
Most Similar (Top) Hit BLAST E Value	4.0E-33	4.0E-33	4:0E-33	4.0E-33	3.0E-33 B	3.0E-33	3.0E-33	2.0E-33 A	2.0E-33 A	2.0E-33 B	2.0E-33 A	2.0E-33	. 2.0E-33	1.0E-33 A		7.0E-34	6.0E-34	6.0E-34	5.0E-34	5.0E-34	4.0E-34	4.0E-34	4.0E-34	1.0E-34	
Expression Signal	1.97	1.14	22	1.48	5.08	3.82	1.73	1.02	2.37	6.1	30.71	2.2	2.2	1.61	1.09	2.31	1.48	1.48	2.53	5.04	1.36	0.92	1.38	14.56	1.32
ORF SEQ ID NO:	12158		12668	14327							14791	14876	14876		14340			10518		14839			13133	11530	13600
Exan SEQ ID NO:	7048	7327		9347	6081	1809		9609	9609		9810		0068	6809	0988			6504	6831	8872			8115	6472	8596
Probe SEQ ID NO:	2066	2353	2474	4358	1073	1074	2382	16	105	4295	4826	4922	4922	6	4368	1419	468	468	1841	4893	1952	2647	3099	1478	3589

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Homo saplans WNT3 precursor (WNT3) mRNA, complete cds	Homo saplens WNT3 precursor (WNT3) mRNA, complete cds	RC2-BT0508-240400-016-h08 BT0508 Homo septens cDNA	hh77b08.y1 NCI_CGAP_GU1 Homo seplens cDNA clone IMAGE:2968787 5'	Homo sapiens prohibitin (PHB) mRNA	nas33808.x1 NCI_CGAP_Kid11 Homo septens cDNA clone IMAGE:3258134 3' similar to TR:075912 075912 DIACYLGLYCEROL KINASE IOTA.;	nsa33a08.x1 NCI_CGAP_Kid11 Homo saciens cDNA clone IMAGE:3258134 3' similar to TR:O75912 O75912 DIACYLGLYCEROL KINASE IOTA :	601809588F1 NIH MGC 18 Hamp septens cDNA clare IMAGE:4040324 61	ah53h03.s1 Soares, testis, NHT Homo saplens cDNA clone 1309397.3'	Homo saplens zinc finger protein 208 (ZNF208), mRNA	Homo saplens hypothetical protein FLJ20420 (FLJ20420), mRNA	H.sapiens immunoglobulin kappa light chain variable region L14	Homo sapiens mRNA for KIAA0406 protein, pertial cds	Homo saplens Ring1 and YY1 binding protein (RYBP), mRNA	Homo sapiens cit.2 kinase (CLK2), propin1, cote1, glucocerebrosidase (GBA), and metaxin genes, complete cds; metaxin pseudogene and glucocerebrosidase pseudogene; and thrombospondin3 (THBS3) gene, partial cds.	601109719F1 NIH MGC 16 Homo septens cDNA clane IMAGE:3350405 5	yu98a07.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:241236 5' similar to contains PTR5 repetitive element :	601125260F1 NIH_MGC_8 Homo septens cDNA clone IMAGE:3345063 5	Homo sapiens phospholipid scramblese 1 gene, complete cds	K6832F Human fetal heart, Lambda ZAP Express Homo sapiens cDNA clone K6832 5' similar to REPETITIVE ELEMENT	A971F Heart Homo saplens cDNA clone A971	Homo sapiens mRNA for Gab2, complete cds	hi86a12.x1 Soares NFL T GBC, S1 Homo sapiens cDNA done IMAGE:2979198 3' similar to SW:TR12_HUMAN Q14888 THYROID RECEPTOR INTERACTING PROTEIN 12;	Homo saplens mRNA for KIAA0895 protein, partial cds	TCBAP2E4328 Pediatric pre-B cell acute lymphoblastic leukemia Baykar-HGSC project≔TCBA Homo sepiens oDNA ckone TCBAP4328	
Top Hit Database Source	Į.	N	EST_HUMAN	EST_HUMAN	<u> </u>	EST_HUMAN	ST HUMAN	EST HUMAN	EST HUMAN	N	1	FZ	ZT TA	Į.	<u>F</u>	EST HUMAN	EST HUMAN	EST HUMAN	TN.	EST HUMAN	ST HUMAN	Į	ST_HUMAN	N N	EST_HUMAN	
Top Hit Acession No.	1.0E-34 AY008397.1	Γ			1190	8.0E-35 BF589637.1 E	8.0E-35 BF589837.1 E			5976	8923389 NT	,	3.2	6912639 NT		4.0E-35 BE257907.1		-		38965.1	1909.1	3018413.1	l _			
Most Similar (Top) Hit BLAST E Value	1.0E-34	1.0E-34	1.0E-34	9.0E-35	8.0E-35	8.0E-35	8.0E-35	8.0E-35	6.0E-35	6.0E-35	6.0E-35	5.0E-35 X63392.1	5.0E-35	5.0E-35	6.0E-35	4.0E-35	4.0E.35	3.0E-35	3.0E-35	2.0E-35 N	2.0E-35 T1	2.0E-35	2.0E-35	2.0E-35	2.0E-35	
Expression Signal	0.93	0.93	4.5	1.41	25.01	2.3	23	3.04	1.6	1.65	0.75	1.63	1.05	1.22	1.3	68.29	19.90	52.74	1.68	121	1.09	2.74	1.92	0.84	0.81	
ORF SEQ ID NO:	13944	13945		13573		11770	14771	14688	11437	12008			12781	12977	14285		11866	11600		10192	11208		12688		13823	
ш <u>ш</u> 2		8955	9343		5286	6694	9694		6385		10032		7668	7959	9277	8406	6775	6542		7713	6171	l	7574		8817	
Probe SEQ ID NO:	3957	3957	4352	3580	224	1699	1699	4717	1388	1925	5063	1671	2711	2939	4285	1408	1783	1544	2269	108	1168	2156	2612	3480	3814	

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	TCBAPZE4328 Pediatric pre-B cell acute tymphoblastic leukemia Baylor-HGSC project=TCBA Homo saplens cDNA cione TCBAP4398	Vol9812.rf Sogres feld live spleen (NEIS Home content of NA state 144.0 E-272225 cm	finite 6 Regional genomic DNA specific chiva library Home services CNA All Colors	finfo16 Regional genomic DNA smartfor DNA library House and the China Ch	12-ST0162-131099-008-412-ST0182-Home series chiva	ILZ-ST0162-131099-006-d12 ST0162 Home seniens cDNA	yd83a01.r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:115752 6 samilar to SP-A44392 A44392 RETROVIDI IS DEI ATEN DOLLON VOCCHERIA	Homo saplens hypothetical protein (LOC61233) mRNA	ht09g01x1 NC_CGAP_Kd13 Homo saplens cDNA clone IMAGE:3146256 3' similar to contains MER29.b3 MER29 repetitive element:	ht09g01.x1 NC!_CGAP_Kd13 Homo sepiens cDNA clone IMAGE:3146256 3' similer to contains MER29.b3 MER29 repetitive element	Homo septens transcription alongshow factor B (SIII) and particle of the Crossist - Care	AV650422 GLC Home sablens aDNA chan (3) CCFENR 2	AV650422 GLC Homo sabiens cDNA clone GL CCFF08 3'	Mus musculus activin receptor interacting protein 1 (Ario1-pending) mRNA	Mus musculus activin receptor interscting protein 1 (Ariot-bending) mRNA	RC3-ST0315-180200-013-f12 ST0315 Homo saciens cDNA	CM1-CT0315-091299-063-407 CT0315 Home saplens cDNA	Homo sapiens C-terminal binding protein 2 (CTBP2) mRNA	Hamo sepiens ninjurin 2 (NIN.12), mRNA	Homo sapiens TCL 6 gene, exon 12	UI-HI-BW 1-any-c-12-0-UI.s1 NCI_CGAP_Sub7 Homo saplens cDNA clone IMAGE:30838423*	Homo sapiens Xq pseudoautosomal region; segment 1/2	601285567F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3607289 6"	Homo sapiens chromosome 21 segment HS21C009	Homo sapiens API5-like 1 (API5L1), mRNA	Homo sapiens API5-like 1 (API5L1), mRNA	PM3-BN0176-100400-001-g04 BN0178 Homo septens cDNA	RETROVIRUS-RELATED POL POLYPROTEIN (CONTAINS: REVERSE TRANSCRIPTASE; ENDONUCLEASE)	601298574F1 NIH_MGC_19 Homo saplens cDNA clone IMAGE:3628386 6'
	Top Hit Database Source	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	1	EST_HUMAN	EST HUMAN	-	EST HUMAN	EST HUMAN	1	1	EST HUMAN	EST_HUMAN	-	1	1	EST_HUMAN	T	EST HUMAN	Į.	1	1	EST HUMAN	SWISSPROT	EST_HUMAN
	Top Hit Acession No.	3E247676.1		-				87947.1 E	7705994	E350127.1 E	3E350127.1 E	883	W650422.1 E		7656905 NT	7656905 NT	W821707.1 E	W857579.1 E	4557498 NT	7705622 NT	1B035346.1 NT		J271735.1 NT		6.0E-36 AL163209.2 NT	5729729 NT	5729729 NT	E010038.1 E		4.1
	Most Similar (Top) Hit BLAST E Vatue	2.0E-35	2.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35	1.0E-35/	1.0E-35/	1.0E-35	1.0E-35	9.0E-36	7.0E-36/	7.0E-36	6.0E-36	6.0E-36/	6.0E-36	6.0E-36/	6.0E-36	6.0E-36	5.0E-36	5.0E-36	4.0E-36	4.0E-38 F	4.0E-36 BE38257
	Expression Signal	0.81	2.63	6.23	623	140.65	140.65	12	2.17	1.37	1.37	124	2.49	2.49	4.67	4.67	1.76	2.3	4.78	1.89	5.02	1.16	10.77	24.78	1.37	2.22	222	2.05	1.38	1.78
<u> </u>	ORF SEQ ID NO:	13824		10119	10120		10791		12555	12786	12767	13100	13121	13122	14276	14277	13883	12898		12045		13567	10219	12751	13542	14621	14622	11243	11474	11684
	Exan SEQ ID NO:	8817	9626	6128	6128	2,196	5766	5916	7439	7652	7852	8086	8106	8106	9290	9290	8880	7880	8081	6943	7325	8561	5204	7636	8537	9828	9628	6206	6413	6802
	Probe SEQ ID NO:	3814	4535	45	45	743	743	888	2471	2694	2694	3070	3090	3090	4298	4298	3879	2860	3044	1957	2351	3554	138	2678	3531	4643	4643	1205	1416	1606

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	2820020 Endme NIH 1/2C 7 H.	R0128226RET NIH 1/2C 41 London Septemble CONTROL 18 18 18 18 18 18 18 18 18 18 18 18 18	601282266F1 NIH MGC 44 Home Called College Col	Homo septems neutredo III enha mare nortisto de	descention and the state of the	organis cardinarcempodus samuleted cyclic nucleotide phosphodiesterase (PDE1A) gene, partial cds	name suprems carcium/cermodulan-stimulated cyclic nucleotide phosphodiesterase (PDE1A) gene, partial cds	none septens KIAA0952 protein (KIAA0952), mRNA	Muse indeceded jurisipalism 1 (Jpt-pending), mKNA	CONTROLL OF THE ANGLE TO FIGURE SERVICES CLIVE CIONS IMAGE:3342708 6	G01300838F1 NIH MCC 34 H	RC1-HT0212-131199-031-05 UT0-312 U	RC1-HT0272-434100-021-407-HT0277-V	602136493F1 NIH MGC 83 Home employee a DNA Alana 114 OF Johanna H	Homo satiens human endonesous retrestration W months 42/2008 6	Homo sapiens chimerin (chimaerin) 2 (CHN3) mDNA	DKFZ0434E0422 rt 434 (synonym: blee3) Home content content blees	one serious in the deservation seeks.		nown gene	790h04 st Sogges fold liver onloan third S st U	EST52010 WATM Home series cDNA chare 52410 circles to the case of	DK-Z0434L2418 r1 434 (swoover: thesa) Home series chive Acre Dk-F2-24 545	DKFZ0434L2418 rf 434 (synonym: htes3) Homo septems cDNA close DKFZ0434 2418	EST373222 MAGE resequences, MAGF Homo septems cDNA	EST373222 MAGE resequences, MAGF Hamo sapiens cDNA	601458531F1 NIH_MGC_88 Homp saplens cDNA clone IMAGE:3882088 6	Homo sepiens mRNA for AML1, complete cds	Homo seplens mRNA for AML1, complete cds	AU131202 NT2RP3 Homo sapiens cDNA clone NT2RP3002166 6'
	Top Hit Database Source	EST HUMAN	EST HIMAN	EST HUMAN	N	5	-	: :		EST HIMAN	EST LIMAN	Т	Т	Т	Т	Т		T HUMAN	Т		EST HIMAN	Т	Т	Т	Т	Т	Τ	П	П		EST_HUMAN /
	Top Hit Acession No.	AW247772.1	Γ	Τ	Γ			2404	10484430 NT	3E259287 1	T	E409310.1	E146523.1	E146523.1	F673761.1	F156962.1	4757979	7.0E-37 AL042800.1		5444607.0	ļ		T	-	Γ				П	2.0E-37 D89790.1 NT	
Mare Challes	(Top) Hit BLAST E Value	4.0E-38	4.0E-36	4.0E-36	3.0E-36	3.05-36	3.05-38	3.05.38	3 OF 36	2.0E-38	2.0E-36	1.0E-36 B	1.0E-36	1.0E-36/B	1.0E-36 B	1.0E-36 A	8.0E-37	7.0E-37 /	7.0E.37 A	7 00 97	6.0E-37 R10039 1	4.0E-37	4.0E-37 N62051.1	3.0E-37	3.0E-37	3.0E-37	3.0E-37	3.0E-37	2.0E-37 D89790.1	2.0E-37 C	2.0E-37
	Expression Signal	4.89	0.98	96.0	2.91	13	13	214	6.39	6.65	17.96	1.87	1.85	1.85	1.5	1.42	0.98	2.66	1.7	;	1.57	2.14	0.91	1.95	1.95	1.2	3.82	0.76	1.7.1	1.71	2.18
	ORF SEQ ID NO:		13311	13312	10725	11524	11525	12329	14345	13130	14768	10833	12176	12177	12234		13313		11776	1477		12447		12056	12057				10472	10473	11104
	SEQ ID NO:	7140			5711	6465	6466		L	8112	9786	5892	7065	7065	7119	8282	8288	6263	6700	6700	8832	7331	10009	6952	6952	7411	7914	9815	5455	6455	6072
	SEQ ID NO:	2161	3275	3275	687	1468	1468	2235	4373	3088	4802	874	2084	2084	2139	3269	3276	1265	1705	1705	4955	2357	5038	1967	1867	2441	2895	4831	88	88	1064

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	AU131202 NTZRP3 Homo septiens cDNA clone NT2RP3002168 6	Homo sapiens chromosome 21 segment HS21C047	Homo septens cytochrome P450, subfamily XXVIIA (steroid 27-hydroxylase, cerebrotendinous	Administration   Description   Circ/Alb mitthe	Homo septembs chromosome 21 seminart MS2(2)84	RC3-CT0347-210400-016-h03 CT0347 Homo sapiens cDNA	Homo septens ribonuclease III (RN3) mRNA, complete cds	QV0-FN0180-280700-318-c10 FN0180 Home sapiens cDNA	Homo sepiens Grb2-essociated binder 2 (KIAA0571), mRNA	602018401F1 NCI_CGAP_Bm87 Homo saplens cDNA clone IMAGE:4163992 5	EST384920 MAGE resequences, MAGL Homo septens cDNA	601455722F1 NIH_MGC_66 Homo saplens cDNA clone IMAGE:3858348 5	EST383908 MAGE resequences, MAGL Homo sapiens cDNA	Homo sapiens RIBIIR gene (partial), exon 8	Homo sapiens RIBIIR gene (partial), exon 8	B.taurus mitochondrial espartate aminotransferase mRNA, complete CDS	B. taurus mitochondrial aspartate aminotransferase mRNA, complete CDS	Homo saplens homeobox protein CDX4 (CDX4) gene, complete ods and flanking repeat regions	Homo sapiens HIRA Interacting protein 4 (dna Like) (HIRIP4), mRNA	SSU72 PROTEIN	SSU72 PROTEIN	601157633F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3504272 6	Homo sapiens chromosome 21 segment HS21C048	Homo sapiens SMT3 (suppressor of mif two 3, yeast) homolog 2 (SMT3H2), mRNA	zw30d01.r1 Soeres ovary tumor NbHOT Hamo saplens cDNA clone IMAGE:770785 6' similar to SW:MA12 RABIT P45701 MANNOSYL-OLIGOSACCHARIDE ALPHA-1.2-MANNOSIDASE:	ZW30d01.rf Soares overy tumor NbHOT Home septens cDNA clone INAGE:770786 6' similar to SW-Ma49 RARIT PA5701 MANNOSY. CI IGOSACCHARIDE AI PHA-1 3 MANNOSITASE	Homo sepiens protein phosphatase 2C alpha 2 mRNA. Complete chs	Homo sapiens keretin 18 (KRT18) mRNA	601177386F1 NIH_MGC_17 Homo septens cDNA clone IMAGE:3532580 5'	601177386F1 NIH_MGC_17 Homo seplens cDNA clone IMAGE:3532580 6"
Top Hit Detabase Source	EST_HUMAN	TN	_ <u></u> _Ę		NT	EST HUMAN	Į.	EST_HUMAN	LN LN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	LN	LZ.	Ę	NT	IN	NT L	SWISSPROT	SWISSPROT	EST_HUMAN	TN	TN	EST HUMAN	EST HIMAN	7	Z	EST_HUMAN	EST_HUMAN
Top Hit Acession No.	2.0E-37 AU131202.1	2.0E-37 AL163247.2	- NO3240	4828885 NT	L163281.2	W862082.1	1.0E-37 AF189011.1	1.0E-37 BF371719.1	11436955 NT	8.0E-38 BF346221.1	7.0E-38 AW972825.1	ŀ	5.0E-38 AW971819.1		5.0E-38 AJZ37740.1	25466.1	25466.1	\F003530.1	7549807 NT	53538	53538	3.0E-38 BE279301.1	L163248.2	2.0E-38 5902097 NT	2.0E-38 AA437353.1	2 0E.38 AA497363 1	F070670 1	4557887 NT	3E296224.1	2.0E-38 BE296224.1
Most Similar (Top) Hit BLAST E Vatue	2.0E-37	2.0E-37	2 0E.47	20F-37	1.0E-37 A	1.0E-37	1.0E-37	1.0E-37	8.0E-38	8.0E-38	7.0E-38/	6.0E-38	5.0E-38	5.0E-38	5.0E-38	4.0E-38 Z25466.1	4.0E-38 Z25466.1	3.0E-38 A	3.0E-38	3.0E-38 P53538	3.0E-38 P53538	3.0E-38	2.0E-38	2.0E-38	2.0E-38	2 OE.38	2 OF 38 A	2.0E-38	2.0E-38	2.0E-38
Expression Signal	2.16	1.45	4 04	0.78	3.59	96.0	1.18	2.02	1.69	1.23	5.28	2.89	1.86	4.11	1.09	3.97	3.97	. 2.4	1.58	1.58	1.58	1.26	1.71	8.04	1.7	ţ	60	9	0.76	0.75
ORF SEQ ID NO:		12004	13808				13863	14751	11240	12617	12212	13001	10757	12478	12478	10200	10201			13772	13773		10127	11408	11865	11888		14421		14910
Exan SEQ ID NO:	6072	6908	8801	9118	7017	8140	8857	9767	6203	7396	6602	7987	5740	7356	7356	5189	5189	7026	8620	8768	8768	9472	5132	6358	6603	BBB	8463	9438	9830	8830
Probe SEQ ID NO:	1064	1923	3798	4123	2834	3124	3855	4783	1202	2426	2119	2969	717	2385	4991	119	119	2043	3613	3765	3765	4482	51	1361	1607	1807	3455	448	4953	4953

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Single Exon Probes Expressed in HBL100 Cells

																<u> </u>	-	1	<u> </u>	w	7						4		-
•	Top Hit Descriptor	zu82b02.rf Soares_tests_NHT Homo sapiens cDNA clone IMAGE:742539 5' similar to contains efement MER19 repetitive element ;	Homo saplens guanine nucleotide binding protein-like 1 (GNL1), mRNA	Homo sapians KIAA0173 gane product (KIAA0173), mRNA	Homo saplens cyclin K (CCNK) gens, exon 7	Homo septens low density lipoprotein receptor-related protein 6 (LRP6) mRNA, and translated products	Homo saplens chromosome 21 segment HS21C003	Homo septens chromosome 21 segment HS21C003	Homo septens hypothetical protein FLJ10600 (FLJ10600), mRNA	Homo sapiens ATPase, H+ transporting, lysosomal (vacuolar proton pump) 1940 (ATPBC) mKNA	Homo sapiens estrogen receptor-binding fragment-associated gene 9 (EBAG9) mRNA	wh53f10.x1 NCI_CGAP_KId11 Homo septens cDNA clone IMAGE:2384491 3' similar to TR:P87890 P87890 POL PROTEIN ;	Homo sapiens chromosome 21 segment HS21C027	Homo sapiens X-linked anhidrotito ectodermal dysplasia protein gene (EDA), excn 2 and flanking repest recions		at36b04.x1 Barstead colon HPLRB7 Homo saplens cDNA clone IMAGE:2374063 3' similar to TR:Q15408 Q15408 NEUTRAL PROTEASE LARGE SUBUNIT; contains LTR7.t1 LTR7 repetitive element;	Chlorocebus aethiops mKNA tor ribosomes protein 54A, complete cas		finfo16 Regional genomic DNA specific cDNA library Homo sapiens cDNA done CK12-1		finifo16 Regional genomic DNA specific CUNA library homo sapiens curva done UN 12-1	801301607F1 NIH MGC 21 Hamo sapiens adire imade: 3030209 0	promme-7.001.r bytumor Homo sapiens culvia o	Homo sapiens homogentisate 1,2-dioxygenase gene, compress cus	PMO-BT0340-211299-003-d02 BT0340 Homo sapiens CUNA	Inv21g02.s1 NCI_CGAP_GCB0 Hamb septens GUNA Globe invAGE:1241150 5 Setting to Contemps 1115.5 THR repetitive element;	Homo sepiens chromosome 21 segment HS21C048	RC4-FN0037-290700-011-210 FN0037 Homo septens CUNA	Homo sapiens KVL.Q.1.1 gene
2	Oatabase Source	EST_HUMAN	T	T,	NT	— <u>E</u>	Ę	M	Ш	11	П	EST_HUMAN	FZ	_ <u>_</u> _		EST_HUMAN	Ę	Ę	EST HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	LN	EST HUMAN	EST_HUMAN	ΕN	EST_HUMAN	Į.
	Top Hit Acessian No.	1.0E-38 AA401570.1 E	36288	7681969 NT	F270831.1	4505016 NT	1.0E-38 AL163203.2 N		8922643 NT	4502312 NT	4758229 NT	AI823404.1	AL163227.2	A FR03528 1	T	AI750154.1	AB015610.1	AL163210.2	AA631949.1		AA631949.1	BE409203.1	AI525119.1	AF000573.1	AW372318.1	AA720574.1	AL163248.2	BF370207.1	AJ006345.1
Most Similar	(Top) Hit BLAST E Vatue	1.0E-38	1.0E-38	1.0E-38	1.0E-38 A	1.0E-38	1.0E-38	1.0E-38	1.0E-38	8.0E-39	8.0E-39	8.0E-39		K 0F.30	200		4.0E-39	4.0E-39	3.0E-39	3.0E-39	3.0E-39			2.0E-39			2.0E-39		1.0E-39
	Expression Signal	1.97	3.28	96'0	2.9	0.72	1.27	1.27	1.06	6.42	1.13	64.1	4.22	70,	26.	6.76	50.63	0.7	18.3	18.3	18.3	18.94	8.24	3.1	89.79	2.58			11.08
	ORF SEQ ID NO:		12041			14186			14436	10133	11417		12130			12853	10584	13514	10121	L	10123					12012	L		11635
	SEQ (D NO:	8083	6839	6955	7394	9185	9180	9190	9456	5136	6368		L	١.		7835	929		6128	L	5129	5904	5919	6025		8015		١.	
	SEQ ID	1076	1953	1970	2423	4182	4197	4197	4466	55	1371	1793	2038	8	788	2916	546	3482	48	8	84	886	8	1015	1498	4620	2558	4279	1482

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Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo sanians (VVI OT4 wasa	Homo seniene DKF7-434D941 modeln (DVE-1042-1044) Chita	EST3840R5 MAGE Recall Concerning to the Concerning Conc	EST364085 MAGE received. WACE Line and	Homo sapiens ()KE7A444D344 models ADKE3D44444	Homo sanians (IDD-Allowed American Control of	Homo septems AE-blinding protein 1 (AEBB1) DNA	Homo saciens AE-binding protein 1 (AEBP1) mBNA	Homo septens tissue inhibitor of metalloproteinase 3 (Soreby fundus dystrophy, pseudoinflammatory) (TIMP3)	Homo serviens fremile X membel respondation 4 /Extract - Distra	Homo senione mRNA (~ K1A 4/3/4 - 1.1.	Homo saniens uhdulilin snedific moteone 49 //	7H15A04 Chromosome 7 Hele & CNA I Thron. Language 1-5) (USP13) mRNA	601288968F1 NIH MGC 8 Home sabless cDNA chare IMAGE: 981919 821	EST70527 I cell hambhane Hamb earline ADMA & and Attin	o etta samuel to	EST 1 0027 1 - Gau rymphoma Homo septens CDNA 5' end similar to similar to zino finger protein family Homo septens chromosome 21 septemble 1004 Cons	#31501 x1 NCI_CGAP_PR28 Homo septems cDNA clone IMAGE:2248873 3' similar to TR:073505 073505 POL PROTEIN:	Homo sapiens X-linked anhidrotito ectodermal dysplasta protein gene (EDA), exon 2 and flanking repeat regions	Homo saplens KIAA0433 protein (KIAAA433) RNA	wh12/07.x1 NCI CGAP Kid11 Home sanians cDNA class 14.4 AE: 228 AE 20 E	9952h08.x1 Soares, testis NHT Homo sepiens CDNA clone IMAGE:180864739	XZ4610 X1 NCL CGAP_Ut4 Homo septens cDNA clone IMAGE:2761098 3' similar to SW 1935_MOUSE P97461 40S RIBOSOMAL PROTEIN SG .	AV731601 HTF Homo saplens cDNA clone HTFAZE05 6	Homo sapiens probasome (prosome, macropain) subunit, apha type, 7 (PSMA7) mRNA, and translated products	Homo saplens protessome (prosome, macropain) subunit, alpha type, 7 (PSMA7) mRNA, and translated products
Top 法	Defabase Source	Ę	¥	EST HUMAN	Т									T HUMAN	EST HUMAN	T	T	LO TOWAN	L HUMAN			T HUMAN	Т	$\Box$	Т		
Too Us A see .	No.	AJ006345.1	7857020	1.0E-39 AW951995.1	AW951995.1	7657020 NT	5803210 NT	4765145 NT	4756145 NT	4507512 NT	4503764 NT	B033070.1	4507848					Τ		F003528.1	82117		2.0E-40 AI223036.1	_		4506188 NT	4506188 NT
Most Similar	BLAST E Vedue	1.0E-39 A				L			9.0E-40	9.0E-40		9.0E-40 A	9.0E-40	8.0E-40	8.0E-40	6.0E-40	00.00	5.0E-40	4.0E-40 AI	4.0E-40 A	4.0E-40	3.0E-40	2.0E-40	2.0E-40	2.0E-40	2.0E-40	2.0E-40
- Gympselm	Signal	11.08	4.37	15.01	15.01	7.93	1.74	11.93	11.93	1.08	1.19	3.4	0.88	96.0	4.61	6.7	. 4	1.57	1.77	2.06	8.89	0.99	3.68	47.86	2.37	6.41	6.41
ORFSEO		11536		14500	14501	14538	10587	11253	11254	11480		13870	14193	13000		12724	12725	12815	11917		14247	13996				11973	11974
	g 2	6479		9515		9552			6214	6420	8701	10048	9214	7986	8825	7614	7814	7495	6828	7032	9257	9007	5377	5807	6781	6882	6882
Probe	SEG ID NO:	1482	1499	4625	4525	4564	551	1215	1215	1423	3697	3866	4370	2968	3823	2854	2854	2529	1838	2050	4264	4011	323	786	1790	1894	1894

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Homo septens adenylyl cyclase-associated protein 2 (CAP2) mRNA	601121567F1 NIH_MGC_20 Hamp septens cDNA clane IMAGE:3345784 5	Homo saplens adenyy cyclase essociated protein 2 (CAP2) mRNA	Homo saplens chromosome 21 segment HS21C080	Homo saplens chromosome 21 segment HS21C080	Hamo saplens plasminogen (PLG) mRNA	nc09a09.s1 NCI_CGAP_Pr1 Hamo sapiens cDNA clane IMAGE:1007608	601460375F1 NIH_MGC_68 Homo saplens cDNA clone IMAGE:3883803 6	bb79a10.y1 NIH_MGC_10 Homo sepiens cDNA clone IMAGE:3048570 5' similar to 118:092168 Q92168 SYNTAXIN 17.;	602068604F1 NIH_MGC_68 Hamo sapiens cDNA clane IMAGE:4067736 6'	602068604F1 NIH_MGC_58 Hamo sapiens cDNA clane IMAGE:4067738 5'	Homo sepiens sorting nextn 3 (SNX3) mRNA	Homo sapiens zinc finger protein 200 (ZNF200) mRNA, and translated products	za36a02.r1 Soares fetal liver spleen 1NFLS Homo sepiens cDNA clone IMAGE:294802 6	wp04h04.x1 NCI_CGAP_Kid11 Hamo sapiens cDNA clone IMAGE:2463895 3*	wp04h04x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2463895 3'	Homo sapiens hypothetical protein (FLJ10996), mRNA	Homo sapiens DSCR5b mRNA, complete cds	Homo saplens Down syndrome candidate region 1 (DSCR1), mRNA	yc03e10.s1 Stratagene lung (#937210) Homo sapiens cDNA clone IMAGE:79626 3	QV0-HT0367-150200-114-509 HT0367 Hamo saprens CUNA	AU118344 HEMBA1 Homo sapiens CUNA cione HEMBATUUDDK3 5	ow45e06.s1 Sogres_parathyroid_tumor_NbHPA Homo sapiens cDNA clone IMAGE:1649794 3' similar to TR:000597 000597 CYTOCHROME C-LIKE POLYPEPTIDE.;contains LTR5.b1 LTR5 repetitive element;	ow45e08.s1 Soares_perethyroid_tumor_NbHPA Homo sepiens cDNA clone IMAGE:1649794 3' similar to TR:000597 000597 CYTOCHROME C-LIKE POLYPEPTIDE, ;contains LTR3.b1 LTR8 repositive element;	Homo sapiens gene for activin receptor type IIB, complete cds	Im98c04.x1 NCI_CGAP_Bm25 Hamo sepiens cDNA clone IMAGE;2163958 3' similar to contains OrK.51	UT RESERVE GROUND IN	Harris september 550 kh contin helween AMI 1 and CBR1 on chromosome 21q22; segment 1/3	
Top Hit Database Source		I HUMAN					T_HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN			T_HUMAN	EST_HUMAN	EST_HUMAN			I. I	T HUMAN	ST_HUMAN	EST_HUMAN	EST_HUMAN	ST HUMAN	Į,		ESI HUMAN		
	토	EST	Ę	Ż	K	Ž	EST	ES	ES	ES	ES	ZNT	N	EST	ES	ES	4 NT	¥	2 NT	EST	EST.		<u>S</u>		z		ŭľ.		=
Top Hit Acession No.	5453592 NT	2.0E-40 BE275932.1	6453592 NT	2.0E-40 AL163280.2	AL163280.2	4505880 NT	AA225989.1	1.0E-40 BF036881.1	1.0E-40 BE018348.1	1.0E-40 BF541030.1	BF541030.1	4507142 NT	4508012 NT	21586.	AIB34364.1	A1934364.1	11431114 NT	6.0E-41 AB037163.1	7657042 NT	5.0E-41 T62628.1	4.0E-41 BE156318.1	AU119344.1	Al027117.1	4.0E-41 Al027117.1	4.0E-41 AB008681.1		A1500406.1	4.0E-41 AJ229041.1	L.TAOSZZOA
Most Similar (Top) Hit BLAST E Value	20E-40	20E-40	20E-40	2.0E-40	2.0E-40 AL	2.0E-40	1.0E-40	1.0E-40	1.0E-40	1.0E-40	1.0E-40 BF	1.06.40	1.0E-40	9.0E-41 W	7.0E-41 AI	7.0E-41 AI	7.0E-41	6.0E-41	6.0E-41	5.0E-41		4.0E-41	4.0E-41 AI						╛
Expression Signal	1.63	13	4.08	1.8	1.8	-	1.42	1.42	1.6	0.89	0.89	122	5.47	1.02	1.8		1.23	2.15		1.57	1.45	1.03	10.1	101					3.89
ORF SEQ ID NO:	12200		13080	14717	14718			12631			12734	L	14448					10350				11117	11432	14433					12855
Exan SEQ ID NO:	7085					L				_					1	L	Ľ	L			1	L	6382	1	1		5 6591		8 7838
Probe SEQ ID NO:	2405	2848	3053	4745	4745	5005	872	2548	2814	2883	2863	3224	4478	3715	818	818	5041	278	2052	1766	390	1081	1385	2007	1305	3	1595	2818	Ŕ

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	History Division (Hose of March 1988)	Home saplens PAD-H19 mRNA for peptidokerninne delminases has il	Homo espiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes,	Hunan rhosomal rodein   23e mDNA Accessive at	EST35818 Fmbno Russk I Home emices - DNA F1 4	Human mRNA for KIAA0207 cana complete and	Gaorilla DNA for ZNEBO gene homologi	Human ribosomal profession 1 22a m RNA Ammulate Ac	Hamp satients son of sevenless (Trescabille) hamples 4 (2004) - Divis	Hamo sablens chramosome 21 segment HS21Ch87	Homo saplens chromosome 21 seriment HS24 Cost	60/445847F1 NIH MCC R5 Home emisse abus Accommoder	01445647F1 NIH MCC 65 Dame carried and 41 mm of constant	Mus musculus tribullo elabe & Tuhos)	Homo saniens hymery myself (DDV) (DDV)	Series Constitution (Constitution Continues Constitution) (Series regions	Mus musculus neura precursor cell expressed, developmentally down-regulated gene 1 (Nedd1), mRNA	Homo sapiens chromosome 21 segment HS21C085	Homo sapiens phosphatidylinositol 4-tdnase 230 (pi4K230) mRNA, complete cds	Homo sapiens phosphatidylinostid 4-kinasa 230 (p4K230) mRNA complete cds	nilar to contains L1.t1 L1	seudoautosomal region; segment 1/2	h/31e11.x1 NC  CGAP_Lu24 Home satiens cDNA clone IMAGE:3175052 31	Homo sapiens SET domain and meriner transposase fusion gene (SETMAR) mRNA	Homo sapiens SET domain and mariner transposase fusion gene (SETMAR) mRNA	Homo saplens MHC class 1 region	Homo sapiens MHC class 1 region	Homo sapiens ribonuclease III (RN3) mRNA, complete cds	H. sapiens PROS-27 mRNA	Homo sapiens SNARE protein kinase SNAK mRNA, camplete ods	Homo sapiens regulatory fector X, 4 (influences HLA class II expression) (RFX4) mRNA	Homo saplens zinc finger protein 177 (ZNF177) mRNA
	Top Hit Database Source	Į.	N <sub>T</sub>	5	Ę	EST HUMAN	Ł	K	¥	<u></u>	F		EST HUMAN	Т						·		EST HUMAN		EST HUMAN									
H	<u>§</u>		Z	2	<u>  z</u>		Z	Z		5032106 NT	호	¥	Ĭ		6678468 NT	Z			ξļ	틸	<u>k</u>	ŭ	¥	ŭ	6730038 NT	5730038 NT	Ż	Z	호	Ä	N	4506496 NT	4508008 NT
	Top Hit Acession No.	4.0E-41 X92885.1	3.0E-41 AB030178.1	3.0E-41 AB026898.1	2.0E-41 U43701.1	2.0E-41 AA331940.1	2.0E-41 D86962.1	2.0E-41 X89631.1	U43701.1		6326	2.0E-41 AL163267.2	1.0E-41 BE869735.1	1.0E-41 BE869735.1	8289	8.0E-42 AF003530.1	. {	1808/00	7.0E-42 AL163285.2	4F012872.1	4F012872.1	6.0E-42 AW238656.1	5.0E-42 AJ271735.1	3E217913.1	6730	6730	4.0E-42 AF055086.1	4.0E-42 AF055068.1	\F189011.1	4.0E-42 X59417.1	\F246219.1	4506	4508
	Most Similar (Top) Hit BLAST E Vatue	4.0E-41	3.0E-41	3.0E-41	2.0E-41	2.0E-41	2.0E-41	2.0E-41	2.0E-41 U	2.0E-41	2.0E-41 AL	2.0E-41	1.0E-41	1.0E-41	1.0E-41	8.0E-42	0 00 40	0.0E-12	7.0E-42	6.0E-42	6.0E-42	6.0E-42	5.0E-42	5.0E-42	5.0E-42	5.0E-42	4.0E-42	4.0E-42	4.0E-42	4.0E-42	4.0E-42	4.0E-42	4.0E-42
	Expression Signal	2.21	1.82	3.05	49.39	1.61	5.84	15.51	16.52	0.8	1.15	1.15	1.18	1.16	15.19	6.2	6	5	1.72	3.13	3.13	2.79	5.47	1.36	8.24	3.56	23.3	23.3	4.34	1.88	0.92	4.17	13.19
	ORF SEQ ID NO:	14004	10983	14183	11586	11998	12255	12306	11586	13751	14458	14459	13170	13171	14409	10508	14030			SORT	11901			10489			10792	10793	11087	14055	14087	14107	14422
	Exan SEQ ID NO:	9017	6950	9201	6527	6904	7136	7183	6527	8752	9478	9478	8149	8149	9424	5497	0052	5025	200	3	808	7204	5202	5473	5520	5521	5767	6767	6058	9065	9100	9122	9441
	Probe SEQ ID NO:	4021	88	4208	1789	1918	2157	2206	2755	3748	4488	4488	3133	3133	4434	460	9267	ě	4840	8101	1819	7227	138	435	483	484	744	744	1049	4071	4108	4127	4451

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Table 4
Single Exon Probes Expressed in HBL100 Cells

Process   Exam   Capt SEC   Expression   Capt SEC   Expression   Capt SEC						_													<u> </u>	<u>بة</u>	_1			! =	<u> </u>	_;	<u></u>	1	<u>-</u>	4	<u>up</u>	무	
December   Corp.   Fig.   Corp.   Fig.   Corp.   Fig.   Corp.   Fig.   Corp.   Fig.   Corp.   Fig.   Corp.   Fig.   Corp.   Fig.   Corp.   Fig.   Corp.   Fig.   Corp.   Fig.   Corp.   Fig.   Corp.   Fig.   Corp.   Fig.   Corp.   Fig.   Corp.   Fig.   Fig.   Corp.   Fig.   Corp.   Fig.   Fig.   Corp.   Fig.   Top Hit Descriptor	b14e10.s1 Strategene lung (#837210) Homo sapiens cDNA clone IMAGE:840810 3' similar to contains THR.t2 THR repetitive element;	RCG-TN0079-110900-024-g07 TN0079 Hamo septens cDNA	AC3-NN0070-270400-011-h10 NN0070 Hamo saplens cDNA	2819283.3prime NIH_MGC_7 Hamo septens cDNA clone IMAGE:2819283 3'	Human endogenous retrovirus pHE:1 (ERV9)	JI-H-BI1-afh-e-04-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2721871 3"	Homo sapiens partial C9 gene for complement component C9, exon 1	Homo sapiens partial C9 gene for complement component C9, exon 1	Home saplens NADH-ubiquinone addoreductase AGGG subunit precursor homolog mRNA, nuclear gene	university illustrational configurations	Flomo sapiens NALIT-Lunquingne axidoreduciase Alges subumit precursor nomang minuk, mudaeri gana encoding mitochandrial protein, complete cds	Homo sapiens rec (LOC51201), mRNA	Homo sapiens major histocompatibility complex, class II, DM alpha (HLA-DMA) mRNA	Homo sapiens origin recognition complex, subunit 6 (yeast homolog)-like (ORCEL) mRNA, and translated	products	Homo sapiens KIAA0255 gene product (KIAA0255), mRNA ·	Homo sapiens Golgi vesicular membrane trafficking protein p18 (BET1) mRNA	Homo saplens chromosome 21 segment HS21C067	Homo sapians chromosome 21 segment HS21C080	RC3-ST0197-161099-012-e03 ST0197 Homo sepiens cDNA	Homo sapiens proteasome inhibitor (Pl31), mRNA	Homo sapiens proteasome inhibitor (Pi31), mRNA	Homo sapiens ryanodine receptor 3 (RYR3) mRNA	AV736824 CB Homo sapiens cDNA clone CBLAKH08 5	AV738824 CB Hamo sapiens cDNA clane CBLAKH08 6"	Homo saplens hypothetical protein FL/20297 (FL/20297), mRNA	Homo sepiens hypothetical protein FLJ20297 (FLJ20297), mRNA	Homo sapiens hypothetical protein FL/20297 (FL/20297), mRNA	2822251.5ptime NIH_MGC_7 Homo saplens cDNA clone IMAGE:2822251 6	ne72d08.s1 NCI_CGAP_Ew1 Homo sapiens cDNA clone IMAGE:809803 similar to gb:1,05095 60S RIBOSOMAL PROTEIN L30 (HUMAN);	AV708201 ADC Homo sepiens cDNA clone ADCACC10 6		
Exam (Note)         ORF SEQ (ID) (ID) (ID) (ID) (ID) (ID) (ID) (ID)	Top Hit Databese Source	ST_HUMAN	ST_HUMAN	ST_HUMAN	ST_HUMAN	Ţ	ST_HUMAN	T	T	_			1			Т	T	Т.	1	T	ST_HUMAN	T	П	П	ST_HUMAN	ST_HUMAN	T	П	T	ST_HUMAN	ST_HUMAN	ST_HUMAN	
Exam         ORF SEQ         Expression (To Signal In Dino)         Most Seq In Dino         Most Seq In Dino         Appression (To Signal In Dino)         Most Seq In Dino         Appression (To Signal In Dino)         Ap													11423219 N	5174458 N		4505524 N	7662027 N	5031610 N			W813617.1			4506758	AV736824.1	4V736824.1		8923276	8923276	AW246442.1	AA491890.1	AV708201.1	
Exen SEQ ID NO: Sign NO: SEQ ID NO: Sign NO: T320 12448 11610 1732 12448 6035 11067 6035 11067 6035 11067 6035 11067 6035 11067 6031 11120 6091 6091 6091 6091 6091 6091 6091 609		3.0E-42	2.0E-42	2.0E-42	2.0E-42	1.0E-42	1.0E-42	1.0E-42	1.0E-42	C7 30 7	1.05	1.0E-42	1.0E-42	1.0E-42		1.0E-42	1.0E-42	1.0E-42	1.0E-42	1.0E-42			1.0E-42			Ŀ						П	
Even ORF SEQ ID ID ID ID ID ID ID ID ID ID ID ID ID	Expression Signal	0.9	261	2.82	4.89	2.19	1.09	1.08	1.08	20.77	08.11	11.85	1.13	1.26		5.85	2.28	0.92	1.08	1.89	0.75	2.84	2.94	5.64	12.63	12.63	5.28	5.28	5.28	8.21	24.66	2.86	
	ORF SEQ ID NO:								L																	L					L		
	SEQ ID	5181	6449	7320	7332	5746	6035	6091	6091	1	74//	7742	6657	7442		7913			L		_	L	<u> </u>			_	L		L		1	1	
	Probe SEQ ID NO:	Ş	1452	2346	2358	724	1025	1084	1084		325	1223	1881	2473		2894	3626	3705	3825	4124	4462	4602	4602	4634	\$	844	8	69	8	3556	1324	2516	

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Top Hit Descriptor	Homo saplens chromosome 21 segment HS21C013	EST96033 Testis I Homo septens cDNA 6' end	AV732578 HTF Homo septens cDNA clone HTFANCO8 6	Homo sapiens X-linked anhidrotitic ectodermal dysplasia protein gene (EDA), exon 2 and flanking repeat	regions	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced	H.sapiens gene encoding La autoentigen	AMI-1-EVI-1=AMI-1-EVI-1 fusion protein (reumanged transfocation) (human, leukemic cell line SKH1, mRNA	William, USON INJ	nkoodos si nci com 111 nomo sapara ocina invadezno i 418	qd61c09.x1 Soares_testle_NHT Homo septems cDNA ctone IMAGE:1733988 3' similer to contains PTR7.t3 PTR7 PTR7 repetitive element;	Homo sapiens Ras-like GTP-binding protein (RABZ7A) gene, exans 1b and 2	Homo sapiens Ras-like GTP-binding protein (RAB27A) gene, exons 1b and 2	Homo saplens chromosome 21 segment HS21C084	602022313F1 NCI_CGAP_Brn67 Hamo sapiens cDNA done IMAGE:4157686 6	qh23g01.x1 Soares_NFL_T_GBC_S1 Hamo sapiens cDNA clone IMAGE:18455623	qh23g01.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1845552 9	RC5-BT0503-081299-011-g12 BT0503 Homo sapiens cDNA	RC5-BT0503-081299-011-g12 BT0503 Hamo saplens cDNA	ye39e01.r1 Soares fetal liver spleen 1NFLS Hamo sapiens cDNA clone IMAGE:124920 5	Homo sapiens LIM domain-containing preferred translocation partner in lipoma (LPP) mRNA	Homo sapiens minisatellite ms32 repeat region	Homo sapiens minisatellite ms32 repeat region	Homo seplens chromosome 21 segment HS210084	Homo sapiens chromosome 21 unknown mRNA	Homo sepiens chromosome 21 unknown mRNA	Homo sapiens KIAA0851 gene (partial), XT3 gene and LZTFL1 gene	Homo sapiens KIAA0851 gene (partial), XT3 gene and LZTFL1 gene	Homo sepiens chromosome 21 segment HS21C103	111d02x1 NCL CGAP Part Homo saplens cDNA clone IMAGE:2130147 3	Homo saplens karyopherin alpha 6 (importin alpha 7) (KPNAG), mKNA	60149152811 NIH MGC G9 Hamp septents active civile invocations of
Top Hit Detabase Source		EST_HUMAN	EST_HUMAN					 		ESI HOMAN	EST_HUMAN		  -		EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN								1		EST_HUMAN		EST HUMAN
	호	ES	SE	_	벌	Z	E	!	Ž	낔	<u> </u>	Ę	Ę	토	ន	S	EB	8	S	ES	N)	뉟	뉟	뉟	Z	호	ĸ	N	Ā	ŭ	픣	<b>"</b>
Top Hit Acession No.	√L163213.2	5.0E-43 AA382780.1	6.0E-43 AV732578.1		AF003528.1	3.0E-43 AF223391.1	3.0E-43 X97869.1		509002.1	3.0E-43 AA548154.1	AI190764.1	AF154836.1	AF154836.1	1.0E-43 AL163284.2	BF348283.1	A1222985.1	AI222985.1	AW373185.1	AW373185.1	R06035.1		_	-		AF231919.1	AF231919.1	AJ289880.1	AJ289880.1	AL163303.2	A1435225.1		BE880626.1
Most Similar (Top) Hit BLAST E Value	5.0E-43 A	5.0E-43	5.0E-43		4.0E-43 A	3.0E-43	3.0E-43	1	3.05-43	3.0E-43	2.0E-43.A	1.0E-43	1.0E-43	1.0E-43	1.0E-43	8.0E-44	8.0E.44	8.0E-44	8.0E-44	7.0E-44	7.0E-44	7.0E-44			7.05-44	7.0E-44	5.0E-44	5.0E-44	4.0E-44	4.0E-44		3.0E-44
Expression Signal	1.96	3.04	1.62		6.71	3.19	4.45		RZ:	0.69	21.1	2.07	2.07	1.71	4.87	4.62	4.62	1.2	1.2	1.08	1.31	2.47		2.74	1.21	121		1.86	3.09		1.97	1.98
ORF SEQ ID NO:		10541			11007		11724			14148		11669					10939	14692			12270	12935				14099			13358			12546
Exam SEQ ID NO:	9029	6535	1		7696	6195			8489	9162	5246	L		<u> </u>		L			L		L	7915	7915	8775	9113	9113					L	7429
Probe SEQ ID NO:	142	499	2773		958	1194	1656		3491	4167	183	1610	1610	1666	2652	879	878	4721	4721	651	2772	2896	2896	3772	4119	4119	် ရှိ	330	3330	4854	1748	2460

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Single Exon Probes Expressed in HBI 100 Cells	Top Hit Descriptor		2018b05.r.1 Strategene fetal retine 037202 Home sentence Chile	Homo saplens DEADIH (Asp-Glu-Ale-AspAHs) how	Homo saplens DEAD/H (Asp-Glu-Ala-Asp-His) town polyagous 1 (DDX1) mRNA	Homo sapiens transmembrane trafficking protein (Tritos) - protein (DUX1) mRNA	Homo sapiens transmembrane trafficking process. (1 Mr 21), mrtva	Homo sepiens RAB36 (RAB36) mRNA, complete cds	PASSES NG CGAP_Lu24 Homo sepiens cDNA clone IMAGE 3182238 3' elmites to cur oxec	TACOS OXIS I EROL-BINDING PROTEIN.	Homo saplens tissue-type bone marrow zinc finger protein 4 mRNA complete 24	Homo sapiens adaptor-related protein complex 4, sigma 1 subunit (Cl ABSA) - Balla	Homo sapiens DNA for amyloid precursor protein, complete eds	PM4-SN0016-120500-003-a04 SN0016 Homo sapiens cDNA	Homo seplens exysterol 7elpha-hydrexylase (CYP39A1) mRNA	Homo sepiens exysteral 7alpha-hydrexylase (CYP39A1) mRNA	Homo saplens Misshapen/NIK-related Knase (MINK) mRNA	Hamo sapiens Misshapen/NIK-related kinase (MINK) mRNA	RC1-CT0249-030300-028-h12 CT0249 Homo saniens CONA	RC1-BN0039-110300-012-b01 BN0039 Homo saniens CONA	Homo sapiens chromosome 21 segment HS21C103	contains THR.t3 THR remetitive december 1999 Homo septems cDNA done IMAGE:773763 5' similar to	2w53d02.rl Soares, total fetus. Nb2HF8_9w Homo sapiens cDNA clone IMAGE:773783 st ===================================		ation 3, JM11 protein, JM4 protein, JM5 protein, T64 protein,	nd Synaptophysin genes,	A done IMAGE:811094 3'	fromo sapiens alpha satellite DNA, M1 monomer type					landard symbol and name) (TFG) mRNA	
Exon Probes	Top Fitt Dertabese Source		EST HUMAN	Ż	N.	Z	Ę	2	FST HIMAN			111	- Is	HUMAN		=			HOMAN	HOMAN		EST_HUMAN	EST HIMAN	7		7	LICHAN							
Single	Top Hit Acessian No.	4 44000000	080144				5803200	AF133588.1	BE465325.1	AF070851 1	5001000	DA7A7E	į	900	1700120	1 N 82 TOO 1 1	N 450 COL	AW/9694994	7	AVV 884603.1	T	AA434564.1 E	AA434654.1 F.	Γ			T	T	111	IN IBOSSO	517A749 AIT	6174718 NT	L160131.1	1
	Most Similar (Top) Hit BLAST E Value	205	77 10 0	200-44	205 44	200	2.0E-44	2.UE-44	2.0E-44		2.0E-44	2.0E-44	2.0E-44		2.0F-44	10F 44	1 OF 44		1 05 44			1.0E-44	1.0E-44 A		1 05-44		1 0F 44 A			D OF AS	8.0E-45	8.0E-45	14	
	Expression Signal	BOB	2 68	2 FB	4.87	4 87	00.7	BRIT	1.58	2.43	3.54	1.97	1.65	1.01	1.9	8.52	8.52	1.72	8	3 88		3.27	3.27		8	4.21	0.76	0.76	1.61	1.61	4.1	8.47	1.91	
	ORF SEQ ID NO:	13050	11071	11072	11228	11227	11337		1398	12186		13424	14411	14623	14624	10130	10131	10606				12259	12280		12758		14900	14901	14418	14419	12538	14862	-	
	SEQ ID NO:	8041	6042	6042		6189	6291		8348	70/2	200	838	9427	8829	9629	5134	5134	5607	6180	6537	3	745	7142		. 7643	8648	9922	8922	9435	9435	7423	6886	9062	
	Probe SEQ ID NO:	3024	1032	1032	1188	1188	1293			3	983	338	#37	\$	4844	8	ន	575	1178	1539	8	3	2,000		2885	3642	4945	4945	4445	4445	2453	4910	2887	
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	Top Hit Descriptor	7481g01 x1 Lupski_dorsal_root_ganglion Homo sapiens cDNA clone IMAGE:3279408 3'	7481g01.x1 Lupski_dorsal_root_ganglion Homo saplens cDNA clone IMAGE:3279408 31	no54e09.s1 NCI_CGAP_SS1 Home sapiens cDNA clone IMAGE:1104520 3' similar to gb:X63741_ma1 FIBULIN-1, ISOFORM A PRECURSOR (HUMAN);	hi86c03.x1 NCI_CGAP_Lu24 Homo sepiens cDNA clone IMAGE:3008836 3' similar to gb:X14008_ms1 LYSOZYME C PRECURSOR (HUMAN);contains element MER37 repetitive element ;	hissco3.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3008836 3' similar to gb:X14008_m41 LYSOZYME C PRECURSOR (HUMAN);contains element MER37 repositive element ;	Human endogenous retrovirus RTVL-H2	Homo sapiens mRNA for KIAA0622 protein, partial ods	Homo sapiens mRNA for KiAA0622 protein, partial cds	Homo sapiens acidic 82 kDa protein mRNA (HSU15552), mRNA	Homo sapiens VAMP-associated 33 kDa protein mRNA, complete cds	Homo saplans mitogen-ectivated protein kinase kinase kinase kinase 3 (MAP4K3), mRNA	H.sapiens Ig lambda light chain variable region gene (7c.11.2) germline; Ig-Light-Lembda; VLembda	H. sapiens ig lambda light chain variable region gene (7c.11.2) germline; ig-Light-Lembda; VLambda	ne08a09.s1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE:890408 3' similar to contains 1 HK.b2 1 HK repetitive element ;	Homo sapiens Bruton's tyrosine kinase (BTK), alpha-D-galactosidase A (GLA), L44-ilke ribosomal protein (L44L) and FTP3 (FTP3) genes, complete ods	z59e02.rl Sogres_testis_NHT Homo sepiens cDNA clone IMAGE:726650 5 elmitar to SW:RSP1_MOUSE. Q01730 RSP-1 PROTEIN.;	Homo sapiens cell division cycle 10 (hamologous to CDC10 of S. cerevisiae) (CDC10) mRNA	Homo saplens KIAA0555 gene product (KIAA0555), mRNA	Homo sapiens KIAA0555 gene product (KIAA0555), mRNA	EST390625 MAGE resequences, MAGP Homo sapiens cDNA	EST48b095 WATM1 Homo saplens cDNA clone 48b085	np76b02.s1 NCL_CGAP_Pr2 Homo sapiens cUNA clone IMAGE:1132355 similes to giz.x1 of 11 n.eapens MT-11 mRNA. (HUMAN);	Homo sapiens mRNA for KIAA0980 protein, partial ods
	Top Hit Datebase Source	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	Ţ	T.	T.	T.	11	Ë	N	_F	EST_HUMAN	T	EST_HUMAN	7	17	F	EST_HUMAN	EST HUMAN	EST_HUMAN	IN
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	Top Hit Acession No.	5.0E-46 BE677194.1	3E677194.1	4.0E-46 AA601143.1	4.0E-46 AW 770544.1	4.0E-46 AW770544.1	4.0E-46 M18048.1	AB014522.1	4.0E-46 AB014522.1	7857	16021	. 4506	3.0E-46 Z73880.1	Z73660.1	2.0E-46 AA468648.1	U78027.1	2.0E-46 AA399288.1				1.0E-46 AW978516.1	1.0E-46 H97330.1	1.0E-46 AA631912.1	1.0E-46 AB023197.1
	Most Similar (Top) Hit BLAST E Value	5.0E-48	5.0E-46	4.0E-48	4.0E-46	4.0E-46	4.0E-48	4.0E-46	4.0E-46	3.0E-46	3.0E-46 AF	3.0E-46	3.0E-46	3.0E-46 Z7	2.0E-46	2.0E-46 U	2.0E-48	1.0E-46	1.0E-46	1.0E-46	1.0E-46			
	Expression Signal	1.07	1.07	2.5	8.01	9.01	2.58	76'0	76.0	1.23	2.18	0.72	1.22	1.22	8.39	2.53	1.31	5.7	1.27			2.62	8.47	3
	ORF SEQ ID NO:	13484	13485		11735	11736	12737	L		12319				14606	10884	<b>1</b>						12431		
	Econ SEQ ID NO:	8458	8458	5663	9661	9861	7825	9285	9285	7199	7335	9262	9615	9615	5847	6597	9798	8212	6536	6536	7195	7310	8182	9708
ŀ	Probe SEQ ID S NO:	3450	3450	835	1885	1885	2888	4283	4293	2222	2361	4269	4630	4630	827	59	4815	1243	1538	1538	2218	2336	3176	4723

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Single Exon Proper Symmetry 1: 1:15:25	LApressed in HBL100 Cells	Top Hit Descriptor		Homo saplens Xq pseudosurbsomal region: serment 475	hi83e04.x1 NCI_CGAP_Lu24 Homo espiens cDNA clone IMAGE-20mes 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	INTEGRAL 12.4 KD PROTEIN.;	mano sapiens HLA-C gene, exon 5, individual 18323	nomo septens HLA-C gene, exxn 8, Individual 18323	Homo seplens protein rivosatudes o	Hamo sapiens 959 kb confin hattern 1111	Homo sapiens chromosome 21 segment 136	Homo sapiens E1A binding partein 2004 (ED2004)	601497639F1 NIH MGC 70 Home sentence - DATA	801497639F1 NIH MGC 70 Home series and 11 11 11 11 11 11 11 11 11 11 11 11 11	1954b04.s1 Soares multiple sciencis 2011-144.s5 Li	Homo sapiens chromosome 21 segment HC24 Code	Homo sapiens glutamate receptor, Innotronic France, 775	Homo saplens nuclear dual-specificity nhose had a control of the c	Human T-cell receptor active aphachain mona 4	Homo saplens myosin phosphatase, famed submitted of American line, complete cds	Homo sapiens chromosome 21 segment H2210no	Homo sapiens chromosome 21 segment HS21 Cana	Homo saplens KIAA0428 gene product (KIAA0428) BNIA		73.			Homo saplens Rev/Rex ectivation domain binding makers 1755	EST377239 MAGE resequences, WAGI Homo saviens child	qp99h03.x1 Soares_fetal_jung_NbHL19W Homo saniens_chwa	601156321F1 NIH_MGC_21 Homo septens CDNA Anna MACCASSASS	GE:3130805 F	Т	spliced spliced and manner appeal E subunit (CACNA1E) gene, exons 7-49, and partial ride when the canal spliced		LAnsa d'array
Exon Probe	王 &	Source		٤	EST HUMAN		Ę		Z	Z.	L	Þ	EST HUMAN	7	T_HUMAN									HOMAN		Т	HOMAN		Т		FOT HUMAN	_	T NOWAN			
Single	Top Hit Acession	g 2	A location	MZ/1/35.1	AW770928.1	Y18536.1	Y18536.1		5453955 NT	1	AL163246.2	92	BEOGRAPA	T	T,		TN 811900+	T		2 OF 47 M 48200	T	2400	2.0E-47 AA524814.4	1000	8	T	0707	2	T			L		223391.1 NT	4501900 NT	
	Most Similar (Top) Hit	Value	0.05.47	100	9.0E-47	8.0E-47	8.0E-47	100	8.05.47	0.0E-47	_	3 05.47							2 OF 47	20F 47	20E-47 A	20E-47	20E-47 A	20E-47	2.0E-47 AA569592 4	2.0E-47 AA5695024	20E-47	2.0F-47 AWORAGO 4	1.0E-47/Al3834294	1.0E-47 BF 280477 4	1.0E-47 RF	1.0E-47 AV		9.0E-48 AF223391.1	8.0E-48	
	Expression		3.81		2.41	80.8	90.6	90	17.5	123	5.93	3.84	3.84	4.98	7.81	0.94	8.7	1.25	129	2.67	2.67	7	3.76	1.6	1.82	1.82	2.23	1.32	4.13	217	2.17	3.23	١	3.80	1.49	
	ORF SEQ ID NO:				14747		11853	12717	1289	12559	11423	10580	10581	10860	10982	13267		14214	10227	11001	11002	11616	11705	14194	14240	14241	14347	14640	11426	13732	13733	14854	11892	3		
	SEQ ID	į	5778		9760	1	\$	7804	7978	7446	6373	5275	5575	6828	5948	8245	8861	8231	5213	6963	5969	6555	9635	9215	9253	8263	9367	9652	6378	8735	8735	8883	8572	6230		
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Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo sepiens eminoscylase 1 (ACY1), mRNA	highbos.x1 NCI_CGAP_Lym12 Hand explores CDNA clane IMAGE:3001133 3" similar to gb:X84707	BREAST BASIC CONSERVED PROTEIN 1 (HUMAN);	h/6/1603.x1 NCI_CGAP_Lym12 Homo sapients GUNA Giare invage3071.53 5 311111	Homo sapiens mRNA for KIAA1209 protein, partial cds	Homo saplens mRNA for KIAA1209 protein, partial cds	Homo contains forsted like kinese 1 (TLK1), mRNA	Tolling depictions SET demails and mariner transposesse fusion perio (SETMAR) mRNA	CORD Sapients OCI Contest to Homo seniers CDNA clone IMAGE 2398613 3"	Widelius XI INC. Cont. Material Administration (PDE1A) mRNA	Homo appears principlications and the companies of CXORF6 mRNA	Homo septens critoring and begins frame 8 (CXORF8) mRNA	Homo sapiens critomosumo A upon teating many of Coord Constitution in MAGE:428844 67	204g03.r1 Soares_leta_liver_spiedn_livrLS_S1 nouls seprens colin constant and spiedness for the constant and spiedness for t	2x80c03.r1 Soares overy tumor Nortici India septients CDIVA care CR17-28	finite? Regional genomic DINA specific duly a library Fortis Backy HCSC projectaTCBA Homo	TCBAP103842 Pediatric pre-8 cell acute lympirtuitaste teurentita payan 1000 pr. 900 to 1000 pr. 1000 pre-8 cell	September County Graduate Home September CONA clane FB2E23'end	PLEAT   Use a September   Home suppers CDNA clone FB2E2 3'end	FEZZZ FEBL Man, Judanycz Francis CONA clone IMAGE: 2689242 3'	Home sanians disnibitin resistance associated overexpressed protein (LOC51747), mRNA		Homo sapiens amyold beta (A4) precured protein (protesse nedn-II, Alzhelmer disesse) (APP), mRNA	Homo saplens EBNA-2 Co-activated (100/D) (P100), illinuta	Homo sapiens EBNA-2 Co-scuvatu (1905) (1905) III. S. S. S. S. S. S. S. S. S. S. S. S. S.	Homo capiens rank birding mount beautiful and the second s	Homo sapiens chromosome at equinal not to re-	Home saplens chromosome 41 administrations and a segment	Human endogenous reuoviral Divo (TT), contentin PD7 domain, complete cds	Mus musculus mysruz miny in injoin comming: ATPase, 4 (PSMC4) mRNA	Homo sapiens protessome (prosonie, literarytes) 263 subunit, ATPase, 4 (PSMC4) mRNA	Home sapiens protested in Courte, Internation 265 subunit, ATPasa, 4 (PSMC4) mRNA	Homo capiens processorie (process) interpretation	
	Top Hit Detabase Source			EST HUMAN	FST HUMAN						EST HUMAN				EST HUMAN	EST HUMAN	EST HUMAN		EST HUMAN	ESI HUMAN	EST HUMAN	FOI HOMAIN		_ <b>-</b> E		E	<u> </u>	PA TA	L	L	Ę	5	5	7	
L		호	L	ES.	25	E			<u>z</u>	z		틸	딁	딁	ES	监	ü			4		<u> </u>	計	88	증	릙	32 N	Z	Z	Z	_	ᇷ	象	8	
	Top Hit Acession No.	TN 001900		8.0E-48 AW 768477.1	9 OF 48 AW788477 1	A DOSSOSE 4	Appeace 4	203303	6912/19 NI	E730038 NT	AI761111.1	4828891 NT		4885170 NT	AA009541.1	AA465007.1	AA631940.1				T03176.1	AW47087	N 48690//	4502166 NT			5032032 NIT	AL163302.2	AL163246.2		AB026497.1			572990 NT	
	Most Similar (Top) Hit BLAST E Value	B OF AR		8.0E-48/	90 00	100	7.05-40	7.0E-48	7.0E-48	7.0E-48	6.0E-48	5.0E-48	3.0E-48	3.0E-48	3.0E-48	2.0E-48	2.0E-48		2.0E-48	2.0E-48	2.0E-48	2.0E-48	1.0E-48	1.0E-48	1.0E-48	1.0E-48	1.0E-48	1.0E-48	1.0E-48	1.0E-48	8.0E-49	7.0E-49	7.0E-49	7.0E-49	
	Expression Signal	1 50	5	3.91		120	1.31	19.03	1.05	6.73	7.99	1.51	28.02	28.02	0.89	1.14	2.18		0.95	1.1	1.1		8.01	4.37											
	ORF SEQ ID NO:	1	1	13092		13053			11523	11854	13537	13272		12016		10071	L	L	14376	14769	14770	15000	10135	10922		L			L					١	
	SEQ ID		250	8079		8079	5524	5524	6464	6593	8526	10046	L	L	L		L		9392	9787		10033	6137	5884	L	Ĺ	L		L	L			L	200	
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Single Exon Probes Expressed in HBL100 Cells

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hit bese Top Hit Descriptor irre	Homo septems professome (homowae mannock) 265 enterus ATO 4 John 2014	Homo satiens and seams (processes marrowell) 262 submit ATD 4 (DOLLA)	Homo saplens protessame (presume, mecroneln) 263 eurhink ATDess 4 (FOWC4) mrtyA	Homo sapiens chromosome 21 segment HS21C084	ba55g05.x1 NIH_MGC_10 Homo sapiens cDNA done IMAGE:2900504 3' similær to gb:X17206 403 RIBOSOMAL PROTEIN S4 (HUMAN); gb:M20632 Mouse LLRep3 protein mRNA from a repetitive element,	Continues (MOCSE);	601457738F1 NIH MGC 68 Home september 2014 Active IMA CE 2001272 5	DKF207614138 s1 761 (supprise beauty) Homes sections CNN states DVF2726444829	Homo sapiens chromosoma 21 segment HS21C010	Homo saplens chromosome 21 segment HS21C010	平28c07.rf Strategene neurospithelium (#837231) Homo sapiens cDNA clone MAGE:610860 6' similar to TR:6233228 G23328 RTVI -H PROTEIN :contains   TE2 rs   TE2   TE2 - All reconstitution of the clone in the clone in the contains of the clone in the clone	Homo saplens putative furnor suppressor ST13 (ST13) mRNA, complete offs	Homo sepiens similar to ribosomal protein S27 (metalliopanstimulin 1) (H. sapiens) (LOC63392). mRNA		mRNA for ecety-CoA carboxylase	ze31c05.r1 Soares retina N2b4HR Homo saptens cDNA done IMAGE:360584 5' similar to contains L1.t3 L1 repetitive element;	Human type IV collagen (COL 4A8) dene. exon 40	Homo sapiens ADP/ATP carrier protein (ANT-2) gene, complete cds	MR3-HT0487-150200-113-g01 HT0487 Homo saplens cDNA	yx23d08.r1 Soares melenocyte 2NbHM Hamo sapiens cDNA clone IMAGE: 282571 67	601458531F1 NIH_MGC_66 Homo saplens cDNA clone IMAGE:3882086 5	Homo sapiens keratin 18 (KRT18) mRNA	601115769F1 NIH_MGC_16 Homo sapiens cDNA clane IMAGE:3356273 5	Homo sapiens chromosome 21 segment HS21C002	Homo saplens mRNA for VIP receptor 2	Hamo saplans mRNA for VIP receptor 2	Homo sapiens actinin, alpha 1 (ACTN1) mRNA
Top Hit Database Source		5	5	N.		EST HIMAN	EST HUMAN	EST HUMAN	NT	N.	EST HUMAN	N	<u>-</u> 1	EST HUMAN	N.	EST HUMAN	¥	N	EST_HUMAN	EST_HUMAN	EST_HUMAN	H	EST_HUMAN	IN	NT	NT	
Top Hit Acession No.	5728990 NT	5729990 NT	5729990 NT	AL 163284.2	2007247W	Ī	T	Γ	Γ	Γ	AA172121.1	J17714.1	11436356 NT	AW189633.1	(68968.1	AA016131.1 E	J46999.1	78810.1	3E165980.1	N26446.1	3F035327.1 E	4557887 NT	E255216.1	L163202.2	95097.2	<b>35097.2</b>	4501890 NT
Most Similar (Top) Hit BLAST E Value	7.0E-49	7.0E-49	7.0E-49	7.0E-49	805.40	6.0E-49	6.0E-49	6.0E-49	5.0E-49	5.0E-49	6.0E-49	5.0E-49	5.0E-49	4.0E-49	3.0E-49)	3.0E-49	3.0E-49	3.0E-49	2.0E-49	2.0E-49	1.0E-49	1.0E-49	1.0E-49 B	8.0E-50 A	8.0E-50	8.0E-50 >	8.0E-50
Expression	3.24	2.89	2.89	3.59	18031	0.80	0.99	0.98	7.25	7.25	3.85	7.57	9.74	22.88	6.0	1.21	2.5	0.94	3.06	1.44	5.66	27.09	6.52	2.76	1.82	1.82	13.5
ORF SEQ ID NO:	10448	10445	10446	11239	10274	11387	11388	13983	li	10742	11836	12750	13236	10658	10591		14790			13187		11675	11843	10244	10749	10750	11789
Exan SEQ ID NO:				6202	6261	8838	6338	9895	5726	6726	6751	7635	8216	5556	6590	7538	8086	9970	5681	8167	5906	6519	6757	5234	5733	6733	6721
Probe SEQ ID NO:	392	393	393	1201	197	1340	1340	3999	702	702	1768	2877	3199	621	556	2575	4824	4999	653	3151	888	1522	1765	169	709	60 20 80 20 80 20 80 20 80 20 80 20 20 20 20 20 20 20 20 20 20 20 20 20	1726

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Hamb saniens p47 // OCS/874/	Hamp serviens 647 // OC\$1674) mBN3	Homo sabions capulan portion (portin florent)	601589565E1 NIH MGC 7 Homosepher PINA Alam 14A CE-201257 EI	CM0-BT0792-300500-398-bd5 BT0792 Home seniens cDNA	CM0-BT0792-300500-398-b05 BT0792 Homo sepiens cDNA	no54e09.s1 NCI_CGAP_SS1 Home sepiens cDNA clone IMAGE:1104520 3' similar to gb:X63741_ma1 FIBULIN-1: ISOFORM & PRECLIES OR HILLIAM.	Homo saciens chromosome 21 sermont HSQ1Cv49	Human endocencus retroduis RTVI H2	601109717F1 NIH MGC 18 Home servine CONA clare IMAGE 232 EASO ET	ODGSTOR STATE HOW SERVING STATE INTO THE PROPERTY OF	hg28e01.x1 NCI_CGAP_GC8 Homo sepiens cDNA clone IMAGE:2948744 3' similar to SW:C1TC HUMAN	P11588 C-1-TETRAHYDROFOLATE SYNTHASE, CYTOPLASMIC; contains Alu repetitive element,	Homo sepiens MHC class 1 region	Homo sapiens midline 1 (Optiz/BBB syndrome) (MID1) mRNA	Homo sapiens decorin D mRNA, complete cds, alternatively soliced	Homo saplens serine palmitoy transferase, subunit II gene, complete cds: and unknown genes			1/2	np98e09.s1 NCI_CGAP_Lu1 Homo saplens cDNA done IMAGE:1142440 3' similar to gb:X12671_ma1 HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN A1 (HUMAN):	xn34e03.x1 NCI_CGAP_Kid11 Homo sepiens cDNA cione IMAGE:2695564 3' straiter to TR:092340 092340 ATYPICAL PKC SPECIEIC BINDING DECTEIN :	QV4-NT0028-200400-180-d05 NT0028 Homo sepilens cDNA	203340 ATYPICAL PKC SPECIEIC BINDING DECTEIN .	DKFZ043482229 rf 434 (synonym: bles3) Home serviens cDNA close DKEZ-434B2229 rf	DKFZp434B2Z29 r1 434 (synonym: htes3) Home septens cDNA clone DKFZp434B2229 r	UI-H-BW0-alp-b-05-0-UI.s1 NCI_CGAP_Sub8 Homo saplens cDNA clone IMAGE:2729817.31	Homo sapiens putative DNA binding protein (MBS), mRNA
Top Hit Detabase Source	  -  2	Ę	Į	EST HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	N.	Į.	EST HUMAN	EST HUMAN		EST_HUMAN	NT	L)	TN	NT	NT	NT	NT	EST_HUMAN	EST HUMAN			Т	Т	Γ	
Top Hit Acession No.	7708394 NT	7708394 NT	4826658 NT	BE794381.1		BF332938.1	AA601143.1		Γ	3E259196.1	AA746142.1			4F055066.1	4557752 NT			386424.1	AL163209.2	U271735.1	AA610842.1	AW274720.1		W274720.1	T			878
Most Similar (Top) Hit BLAST E Value	8.0E-50	8.0E-50	8.0E-50	6.0E-50	5.0E-50	6.0E-50	4.0E-50	4.0E-50	3.0E-50	3.0E-50	3.0E-50		3.0E-50,	Z.UE-50 /	2.0E-50	2.0E-50 /	2.0E-50	2.0E-50	1.0E-50/	1.0E-50 /	8.0E-51	7.0E-51	_	7.05-51	7.0E-51	7.0E-51 /	7.0E-51	6.0E-51
Expression Signal	128	1.29	3.51	0.88	1.19	1.19	1.83	98.0	2.31	1.05	0.89		40.	14.02	5.6	1.29	0.78	1.11	1.74	7.62	12.15	.33	1.83	0.82	1.25	1.25	2.54	1.18
ORF SEQ ID NO:	12500	12501				11834		13393		12532	13262		148/4		11103	11475	13250	14116	10507		14410	12986	13246	13319	14029	14030	14198	11549
Exan SEQ ID NO:	7382	7382	1587	9210	6750	8750	5923	8373	6886	7417	8240		SUOUT POOLS	0100	6071	8414	8228	9133	5496	7279	9425	7969	8224	8294	603	8039	9219	6494
Probe SEQ ID NO:	2411	2411	2627	4217	1757	1757	908	3385	1898	2447	3225		2802	3	200	1417	3214	4138	<u>8</u>	2304	4435	2850	3209	3282	4043	4043	4225	1496

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Homo sepiens KIAA0929 protein Mas2 interacting nuclear tenget (MINT) homology (KIAA0929)	Homo earliens KIAA0929 onthein Mar2 Interaction unclear terms (Alline)	Homo sapiens chromosoms 21 segment HS21C2003	Homo sapiens T-cell tymothoma investion and metastaste 1 (TIAM4) mDNA	Novel human gene mapping to chomosome X	Homo sepiens 26S protessome associated heard homology (POHA) DNA	Homo saciens mRNA for nucleonodo 155	Human Ku (b70080) subunit mRNA complete ode	Human Ku (b70/b80) subjumit mBNA complete ala	Homo sapiens mRNA for KIAA1411 routien medial ade	#61009x1 NCI_CGAP_Pen1 Homo septems CDNA clone IMAGE:2224720 3' similar to gb:M26326	NERALIN, 17PE I CYTOSKELETAL 18 (HUMAN);	181008X1 NCI_CGAP_Pan1 Homo sepiens cDNA clone IMAGE:2224720 3' similar to gb:M28326 KERATIN, TYPE I CYTOSKEI FTAI 18 /HIMANY	2087001.s1 Stratagene hNT neumon (#337233) Home contame ADNA Attack (14) ST-0 10000 at	Novel human gene manning to chomosome 22	Homo sepiens ubiquitin protein ligase E3A (human papilioma virus E6-associated protein, Angelman.	201285804E1 NIEL 18CC 44 Vone Anniel Pris III	601285694F1 NIH MAC 44 Home emission of NA ALCO NA ALC	### ### ##############################	#27g03.x1 NCI CGAP Kid14 Homo septems china che in A CE 2424723 21	Homo saplens eukarvotic translation initiation factor 4A iscform 1 (FIEA41) mBNA	AV742248 CB Homo saplans cDNA clone CRFRCC12 5	nw21g02.st NCI_CGAP_GC80 Hamo septens cDNA done IMAGE:1241138 3' similar to contains THR ts	THR repetitive element;	H.saplens mRNA for laminin-5, alpha3b chath	Homo saplens twodheffcal protein F. 113556 similar in Numer drametreen see John 1919 1905 1905		T -
Top Hit Database Source	<b>F</b> V	<del>_</del>	¥	Ę	Į.	5	Ę	Z.	Į		200	ESI HUMAN	EST HUMAN	Γ	Г		T HI IMAN	Т	Т	Т		T HUMAN	Т	LHUMAN	NT.			
Top Hit Acession No.	7657288 NT	7657288 NT	6320	7500	33204.1	5031980 NT	5.0E-51 AJ007558.1		938.1	37832.1	Γ	1	3.0E-51 AI587348.1 E		Γ	80770	2.0E-51 BE391083 1	T		2.0E-51 Al492415.1	3528	42248.1		8.0E-52 AA720574.1 E		11968028 NT	11968028 NT	
Most Similar (Top) Hit BLAST E Value	6.0E-61	6.0E-51	5.0E-51 AL1	5.0E-51	5.0E-51 AL	5.0E-51	5.0E-51	5.0E-51 M30	6.0E-51 M30	5.0E-51 ABO	2 OF 64	10-20-5	3.0E-51	3.0E-51	3.0E-51	2 0F-84	2.0E-51	2.0E-51	2.0E-51	2.0E-51	1.0E-51	1.0E-51 AV7		8.0E-52 A	8.0E-52	8.0E-62	8.0E-52	
Expression Signal	3.34	16.79	6.86	1.74	1.07	1.42	10.9	3.67	3.67	1.64	20.61		35.84	1.08	2.01	2.45	1.65	1.65	6.36	2.94	45.03	51.24		7.75	1.35	2.31	231	
ORF SEQ ID NO:	12019	13427	١.		11024	11630	12601	13850	13851	14848	10217		11195	11960	14175	10427	10712	10713	11718	13860	10195			10230	11522	11674	11675	
SEQ (D NO:	6920	8401	5804	5815	7738	6568	7485	8843	8843	9879	5201		6161	6871	9193	5414	5704	5704	6645	8654	5186	6428		6217	9463	6610	9810	
Probe SEQ ID NO:	1934	3383	<b>28</b>	क्र	976	1571	2517	3841	3841	4900	135		1167	1882	4200	365	679	679	1649	3648	114	1462	•	151	1488	1614	1614	

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Tap Hit Descriptor	Homo sapiens hypothetical protein FLJ13559 similar to N-mvc downstream manished 3 /FI 1138551 mRNA	Homo septens hypothetical protein FLJ13556 similar to N-mvo downstream remistred 3 / FL 143559 PNA	Homo septens S164 gene, partial cds; PS1 and hypothetical protein genes, complete ods; and S171 gene, partial cds.	H.saplens flow-carted chromosome 6 Hindill freament SCena 1847	Homo sapiens SH3-containing protein SH3GLB1 mRNA, complete eds	Homo saptens nucleoporin (55kD (NUP155) mRNA	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	Homo sapiens 5,10-methyleneistrahydrofolata dehydrogenase, 5,10-methyleneistrahydrofolate cyclohydrolase, 10-formylietrahydrofolate synthetisse (ATTHED) mRNA	Homo seplens hypothetical protein FLJ10675 (FLJ10675) mRNA	Human endocenous retrovirsi DNA (4-1) complete retrovirsi	Human endogenous retroviral DNA (4-1). complete retroviral segment	Homo sepiens KIAA0439 mRNA, partial cds	Homo sapiens mRNA for KIAA1249 protein, partial cds	bb86b07.y1 NIH_MGC_9 Home septens cDNA clone IMAGE:3030421 6' similar to gb:X16493 M.muscutus mRNA for Zof-1 zinc finger protein (MOUSE).	602084710F1 NIH_MGC 83 Homo saplens cDNA clone IMAGE:4248801 5'	Novel human gane mapping to chromosome 20, similar to membrane transporters	qa56e05.s1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAGE:1690784.3*	qa56e05.s1 Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGE:1690784 3'	zu75h12.s1 Soares_testis_NHT Homo sepiens cDNA clone IMAGE:743879 3'	Homo sapiens glutamate-ammonia ligase (glutamine synthase) (GLUL) mRNA	Homo sapiens anysulfatase D (ARSD), transcript variant 1, mRNA	pol-reverse transcriptase homolog (retroviral element) [human, endogenous retroviral element RTVL-Hp1, Genomic, 660 nt]	Homo septens protein kinase, cAMP-decendent, requisitory, type II. beta (PRKAR28) mRNA	Homo sapiens core binding factor alpha1 subunit (CBFA1) gene, exon 3	Homo sapiens mRNA for KiAA1504 protein, partial cds	Homo sapiens heterogeneous nuclear ribonucleoprotein C (C1/C2) (HNRPC) mRNA	Homo sapiens chromosome 21 segment HS21C085	Hamo saplens chromosome 21 segment HS21C085
Top Hit Database Source	_ <del>_</del>	— <b>5</b>	<u>-</u> - <u>-</u> -	Z	N	N	NT	— <b>Ļ</b>	NT.	¥	¥	F	N	EST HUMAN	EST HUMAN	N.	EST_HUMAN	EST_HUMAN	EST_HUMAN	٨T	ΛŦ	_N	5	N-	L	ZT.	NT .	NT
Top Hit Acession No.	11988028 NT	11968028 NT		Γ		4758843 NT	4507500 NT	6174590 NT	11437042 NT	2.0E-52 M10976.1	Γ	2.0E-52 AB007899.1			Γ				1634445.1	4504026 NT	4502238 NT	S61070.1	4506064 NT		9.0E-53 AB040937.1	4758543 NT	4.0E-53 AL163285.2	
Most Similar (Top) Hit BLAST E Value	8.0E-52	8.0E-62				4.0E-52	4.0E-52	· 4.0E-52	3.0E-52	2.0E-52			2.0E-52	2.0E-52	2.0E-52	2.0E-52	2.0E-52	2.0E-52	1.0E-52 A	1.0E-52	1.0E-52	1.0E-62	9.0E-63	9.0E-53	9.0E-53	5.0E-53	4.0E-53	4.0E-53
Expression Signal	6.2	. 6.2	3.39	2.8	1.32	2.08	0.81	1.26	10.25	1.85	1.85	1.15	1	3.12	19.48	3.17	1.1	1.1	1.37	8.69	1.67	1.99	1.3	122	1.19	15.99	1.53	1.53
ORF SEQ ID NO:	11674	11675	11723		11685	11823	13835	14484		10592	10593	11790	12052	12518		14782	14808	14809	10568	11401		13015	13708	14250	14975	13965	10125	10126
Exan SEQ ID NO:	6610	6610	6651		6619	6744	8828	9504	8973	<b>6</b> 592	<b>6</b> 592	6713	6948	7397	7821	9802	9834	9834	5564	8351	7434	8008	8706	8260	10004		5131	5131
Probe SEQ ID NO:	3888	3888	1655	4318	1622	1750	3826	4514	3975	558	558	1718	1964	2428	2882	4818	4853	4853	629	1354	2465	2985	3702	4287	5033	3982	જ	8

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo sapiens hookt protein (HOOK1) mRNA	Homo sapiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes,	2007 V Same Distance at Miles	12 1 Mings 240300 are the last in the control Home explans cDNA clone IMAGE:2558796 3'	EST77525 Parces time III Home Splens cDNA	Homo sapiens Bruton's tyrosine kinase (BTK), alpha-D-galactosidase A (GLA), L44-like ribosomal protein (L44L) and FTP3 (FTP3) genes, complete orls	Homo sapiens ATPase, H+ transporting, lysosomal (vacuolar proton pump) 31kD; Vacuolar proton-ATPase, subunit E. V-ATPase, anthrette (ATPase),	Homo seniena leurine eminonentidoso // OCEAGES - FILLS	dams enfants dilutions della constanti della c	Human Knieonel-raleidd DNA bhraham achdi (7527)	Homo seniese SKADES hample (CKAD Light) Battle Cds	Homo sapiens Xo pseudosimal malor: accompany and accompany and accompany and accompany and accompany and accompany and accompany accompany and accompany acc	Homo explens DNA, DLEC1 to ORCTLA gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes,	60117672FF NIH MCC 17 Umm - This is the second of the seco	EST369619 MAGE recembers. MACE 11	601272863E1 NIH MGC 20 Home environ Child All March 11/1 CE 201 1201 11	Homo sapiens insulin-like growth factor 2 recently (ICEOD) DNA	Homo sapiens ublandin specific prolesse 13 (soperitiess 7.3) (1954) - Data	Homo sapiens ubliquitin specific protease 13 (Isonentkase 1.3) (ISD13) mPNA	contains MER30.43 MER30	NA for monocyte chemotectic protein-2	formo sapiens cDNA clone IMAGE:257399 s'	A CONTRACTOR OF A CONTRACTOR O		(F7n494Mings) mbNA			
	Top Hit Database Source	Z I	<u></u>	FST HIMAN	EST HIMAN	EST HUMAN	_F	_ <u>_</u>	Į.	Ę	Ę				T HUMAN	T	Т				EST HUMAN	Т	EST HUMAN	Т	П				
	Top Hit Acession No.	7705414 NT	AB028898.1	AW050836 1	AW803563 1	4A366556.1	J78027.1	4502316 NT	7705887 NT	1F08382	161873.1	4506962	1.0E-53 AJ271738.1		Ī	Γ	8.0E-54 BE386785.1	610	4507848 NT	4507848 NT		7.0E-84 Y16645.1			3003618.1	8922148 NT	8922148 NT	8922148 NT	4502872 NT
	Most Similar (Top) Hit BLAST E Value	4.0E-53	3.0E-63	3.0E-53		l.	2.0E-63	2.0E-53	2.0E-53	2.0E-53/	2.0E-53 N	2.0E-53	1.0E-53 /	1.0E-53	1.0E-53	1.0E-53 /	8.0E-54	8.0E-54	8.0E-54	8.0E-54	7.0E-54	7.0E-54	7.0E-54 N27177.1	7.0E-54 AI	6.0E-54	6.0E-54	6.0E-54	6.0E-54	8.0E-54
	Expression Signal	98'0	1.47	1.94	0.73	3.58	20.13	7.48	0.92	2.53	2.5	0.92	1.56	1.23	1.54	0.97	4.09	2.71	0.71	0.71	1.58	1.37	424	1.08	5.96	1.73	1.73	21	1.11
	ORF SEQ ID NO:	14843	12667	13659			12360			13210	H	14336	11477	13355	14773	14979	10283	11880	14573	14574	10475	11875	12237	14983	10088	10476	10477	13247	13896
	SEQ ID NO:	9860			9447		7243	- 1		8188	8945	9326	6417	8335	9791	10010	5271	6790	9584	9584	5458	6785	7121	10014	5103	5459	5459	8225	8898
	Probe SEQ ID NO:	4675	2589	3847	4457	455	2266	2466	3147	3172	3947	4365	1420	3325	4807	603	207	1799	4596	4596	383	1794	2142	5043	ಜ	384	384	3210	3898

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Human mRNA for KIAA0077 gene, pertial cds wd26d11.x1 Soares\_NFL\_T\_GBC\_S1 Homo sapiens cDNA clone IMAGE:2329269 3' similar to TR:002711 002711 PRO-POL-DUTPASE POLYPROTEIN; Homo saplens chromosome 21 segment HS21C010
wy60b12x1 Soares\_NSF\_F8\_9W\_OT\_PA\_P\_S1 Homo saplens cDNA clone IMAGE:2552927 3' similar to
TR:Q62084 Q62084 PHOSPHOLIPASE C NEIGHBORING; Homo saptens killer cell lectin-like receptor subfamily G, member 1 (KLRG1), mRNA Homo saptens nuclear antigen Sp100 (SP100) mRNA nt78a09.s1 NCI\_CGAP\_Pr3 Homo saptens cDNA clone IMAGE:1204600 similar to contains element L1 128e04.r1 Soares fetal liver spleen 1NFLS Homo saptens cDNA clone IMAGE:127998 6 similar to n/45g99.s.1 NCI\_CGAP\_Pr9 Homo sepiens cDNA clone IMAGE:998488 similar to gb:X63777 60S RIBOSOMAL PROTEIN L23 (HUMAN); Bu92g03.y1 Schneider fetal brain 00004 Homo saplens cDNA clone IMAGE:2783764 5' similar to EST177696 Jurket T-cells VI Homo sapiens cDNA 6' end similar to glyceraldehyde-3-phosphats 295b09.s1 Soares\_fetal\_liver\_spleen\_1NFLS\_S1 Homo sapiens cDNA done IMAGE:462617 37 295b09.s1 Soares\_fetal\_liver\_spleen\_1NFLS\_S1 Homo sapiens cDNA done IMAGE:462617 37 UI-H-B11-afy-g-09-0-UI.s1 NCI\_CGAP\_Sub3 Homo sapiens cDNA clone IMAGE:2723636 31 Homo sapiens phosphetidylinositol 4-kinase, catalytic, alpha polypoptide (PIK4CA) mRNA Homo sapiens syncylin precursor, mRNA, complets ods Homo sapiens SKAP55 homologue (SKAP-HOM) mRNA 80189230F1 NIH\_MGC\_19 Homo sapiens cDNA clone IMAGE:4128535 6\* EST165371 Colon cardnoma (HCC) cell line Homo septens cDNA 5' end Homo sapiens chaperonin containing T-complex subunit 8 (CCT6) mRNA H. sapiens shc pseudogene, p68 boform ZINC FINGER PROTEIN 84 (ZINC FINGER PROTEIN HPF2) Top Hit Descriptor SW:CUL1\_HUMAN Q13616 CULLIN HOMOLOG 1; IL-BT189-180389-007 BT189 Homo saplens cDNA Homo saplens RFB30 gene for RING finger protein upala belangeri beta-actin mRNA, partial ods Human mRNA for KIAA0077 gene, partial cds SP:C561\_BOVIN P10897 CYTOCHROME H.saplens she pseudogene, p68 laoform Single Exon Probes Expressed in HBL100 Cells epetitive element: dehydrogenase NT SWISSPROT Top Hit Detabase Source EST\_HUMAN EST\_HUMAN **EST HUMAN** EST\_HUMAN EST\_HUMAN **EST HUMAN EST HUMAN** EST\_HUMAN EST HUMAN EST\_HUMAN EST\_HUMAN **EST\_HUMAN EST HUMAN** 뉟 눋 4507164 NT 눚 4502642 NT 4505806 Top Hit Acession 5031900 2.0E-54 AW057524.1 4.0E-54 AA306784.1 4.0E-54 D38521.1 3.0E-54 AA313487.1 3.0E-54 AI908757.1 2.0E-54 AW163175.1 AF110103.1 2.0E-54 AA665008.1 AL163210.2 BF315418.1 6.0E-65 AW 206021.1 Š 4.0E-54 AI835088.1 2.0E-64|AA532925.1 2.0E-54 AF208161.1 6.0E-55 AA704971.1 6.0E-55 AA704971.1 6.0E-54 Y09846.1 6.0E-54 Y09846.1 8.0E-55 Y07829.2 4.0E-54 D38521 7.0E-55 R09346.1 6.0E-54 P51523 4.0E-54/ 2.0E-64 .0E-54 2.0E-54 2.0E-54 2.0E-54 Most Simila BLASTE 堂(金) Value 2.18 140.55 236 3.25 2.65 1.39 30.78 263.62 2.55 <del>1</del>9 1.04 6.13 1.47 1.65 <u>7</u> 7.32 0.92 1.85 <u>8</u> 2.6 1.31 Expression Signal 14664 12185 ORF SEQ 10991 11849 10179 10668 11569 12810 12865 11106 11395 12554 14992 11804 11805 14600 ÖΝΩ SEQ ID 7071 8709 9709 5245 88 5958 5169 7516 5664 6344 6513 7490 7845 6761 8761 8147 7437 8076 10023 <u>8</u> 8728 6728 ÿ 9332 9539 9311 88 Probe SEQ ID 4724 4724 28 82 28 82 1769 1769 182 3131 2 92 සි 1515 2469 3472 1347 4319 2551 2824 4082 5052 1298 1065 1733 2524 1733 ÿ

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Ten Hit Describber		EST370064 MAGE resequences, MAGE Homo seplens cDNA	Homo sepiens RNA binding motif protein, Y chromosome, family 1, member A1 (RBMY1A1) mRNA	Homo sapiens predicted osteoblast probin (GS3788), mRNA	Homo sapiens predicted osteoblast protein (GS3786), mRNA	752b10x1 Soares_NSF_F8_9W_OT_PA_P_S1 Home septems cDNA clone IMAGE:3390043 3' similar to	CONTAINS LT. IX LT REPORTIVE EXEMPTIT;	Trustic separate procedure (procedure, rescupent) establist, suprim type, z (Townyz) mitury.	Torrid equation of control three control management against type, a (1 cm/s) minute	Homo septens discussing forms (such (DCNC) (DCNC)	Homo contant under the control of th	James equients advancement 24 comments INCA CADA	Tromo saprens du concisione 21 segment no 210 un	RCZ-U10023-280700-011-103 U10023 Hamo septens CDNA	Human endogenous retrovirus pHE.1 (ERV9)	Human endogenous retroviral DNA (4-1), complete retroviral segment	Homo sapiens syntaxin-binding protein 1 (STXBP1) mRNA, and translated products	Homo sapiens ubiquitin protein ligase E3A (human papilloma virus E6-associated protein, Angelman syndrome) (UBE3A) mRNA	CM1-HT0876-150800-357-403 HT0876 Homo saciens cDNA	Homo sapiens mannose-6-phosphate receptor (cation dependent) (M6PR) mRNA	Oryctolacus cuniculus New Zeelend white elongation factor 1 alpha (Rabetlaz) mRNA, complete cds	ov85g09.x1 Soares_testis_NHT Homo sepiens cDNA clone IMAGE:1844160 3*	Homo sapiens mRNA for KIAA0903 protein, partial cds	601120116F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:2867027 6'	601120116F1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:2867027 5	Homo saplens SMA3 (SMA3), mRNA	Human mRNA for HLA-A11E, a MHC class I molecute (major histocompatibility complex)	Homo sapiens mRNA for KIAA0406 protein, partial cds	Homo saplens mRNA for KIAA0406 protein, partial cds	Homo sapiens CLP mRNA, partial ods	Homo sapiens mRNA for KIAA1219 protein, partial cds	Homo sapiens 5,10-methylanetetrahydrofolate dehydrogenase, 5,10-methylenetetrahydrofolate ordohydrofase (MTHFD) mRNA	
Top 任用	Source	EST_HUMAN	N	Į.	N		EST HUMAN		1		2 5	F1	- 2	EST HUMAN	NT	NT	M	— <u></u>	EST HUMAN	LZ	<u> </u>	EST HUMAN	F	EST_HUMAN	EST_HUMAN	Ę	TN	F	NT	NT	NT	- <u>1</u>	
Top Hit Acession	ć Ž	\W857894.1	4826973 NT	7661713 NT	7661713 NT		3F081411.1	TIN CORPORA	4300100	4503314 NIT	4507704 NT	5		3E698671.1	<b>K57147.1</b>	M10976.1	4507296 NT	1007708A	3F719986 1	4505060 NT	109823 4	AI028718.1	AB020710.1	BE277861.1	BE277861.1	5803174 NT	X13111.1	AB007868.2	AB007866.2	L54057.1	AB033045.1	F174590 NT	
Most Similar (Top) Hit	BLAST E Value	4.0E-85	4.0E-65	4.0E-55	4.0E-55		4.0H-05	4.0E-00	4.05-30	4.05-05	4.05-55	10.10.1	4.0E-65	4.0E-55	2.0E-55	2.0E-55	2.0E-55	2.05-45	2 OF-65	1.0E-55	4 DF-85	1.0E-55	1.0E-55	1.0E-55	1.0E-55	1.0E-55	1.0E-55	1.0E-55	1.0E-55	1.0E-55	1.0E-55	4 0F.45	
Expression	Signal	6.49	4.62	1.58	1.58		£ 2	CR	CR.I	500	0.0		1:2	2.08	2	1.08	4.78	44	79.67	2.88	113.02	14.24	88.8	9	9	6:36	63.73	3.06	3.06	14.65	0.98	1.83	22.
ORF SEQ	Ö Q	10134	10699	11471	11472		1000			12121		1004			10438		10673	43030									12531	L					
E Gan	ë	7712	2690	6412	6412	1	8480	A S	ROS F	7015	2007	100			5423	5581	6999	7008		L		L			6897	L	L				Ŀ	١.	1830
Probe	ÿ	8	684	1414	1414		1483	4/81	18/8	2032	2032	1477	3206	4985	378	547	841	2880	4820	95	180	569	1132	1911	1911	2263	2446	2483	2483	2538	2717		ROSS

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Table 4
Single Exon Probes Expressed in HBL100 Cells

			T	T	Т	Т	11.	•	Т	Т	Т	7	Т	·		_	Г	, g	<u>ر ر</u>	-	1	4	Ц	5	_	Д	Ц		4	1	1	5,	5,1
Top Hit Descriptor	Homo sapiens 5,10-methylanetatrahydrofolate dehydrogenase, 6,10-methylanetetrahydrofolate	sycianyanderse, 10-formylietrathydroficiale synthetase (MTHFD) mRNA	Homo sapiens chromosome 21 segment HSZ1C067	namo sapiens chromosome 21 segment HS21C010	W448U3.71 Soares fetal liver spleen 1NFLS Hamo sepiens cDNA clone IMAGE:245620 5	Homo sepiens hypothetical protein FLJ20128 (FLJ20128), mRNA	NC3-5 1003-150200-031-811 BT0605 Homo sapiens cDNA YR62g03-11 Soares adult brain N2b5HB557 Homo sapiens cDNA done IMAGE:173044 6' similar to contains	I FIX repetitive element;	RCS-SNUV53-170200-011-h01 BN0053 Homo saplens cDNA	nomo sapiens beta-tubulin mRNA, complete cds	riomo espiens beta-tubulin mRNA, complete cds	Homo sapiens tubulin, beta polypeptide (TUBB) mRNA	Homo sapiens tubulin, beta polypeptide (TUBB) mRNA	Homo sepiens X-linked enhidrotic ectodermal dyspiasia protein gene (EDA), exon 2 and flanking repeat	John Marie Contains have the state of the st	Home series F. 9. 11.	Homo equience of automatice as a 2 (XXN2), mRNA	ST28880 Contelline II U.	EST28880 Cereballim II Hamo sepiens cDNA 6' end	Homo contains MUC Aleas 4	MARCHANIA MOC AA U.	Homo sapiens Down syndrome candidate 1/100001	Homo saplens chromosome 21 segment H321Ch8a	Homo saplans surperkiller drainfile arthaby 2 committees in the saplans surperkiller drainfile arthaby 2 committees in the saplans surperkiller drainfile arthaby 2 committees in the saplans surperkiller drainfile arthaby 2 committees in the saplans surperkiller drainfile arthaby 2 committees in the saplans surperkiller drainfile arthaby 2 committees in the saplans surperkiller drainfile arthaby 2 committees in the saplans surperkiller drainfile arthaby 2 committees in the saplans surperkiller drainfile arthaby 3 committees in the saplans surperkiller drainfile arth	Homo sapiens phosohodidylinosital transfer protein hete (PITONID) - Days	Home sapens checkled transfer process, one (TITLE) INC.	052808.51 Stratanena neuroenithalium (#037231) U.	RC4-BT0310-110300-018-4-0 pT0340 U	RC4-BT0310-110300-015-f10 BT0310 Home confers CDNA	Human GMD phoenhodischesses although the selection of many	Human GMP phosphodiastages alpha author (COP A) TRIANA, complete cas	Homo sapiens mRNA for KIAA1414 protein partiel Att	one manual transactions
Top Hit Database Source	5		Ž	١	NAME OF THE PERSON OF THE PERS	NI Control	NAMOL I COL	EST HUMAN	NAMOR I SE	1	1	= !	5	_ \				T HI IMAN	HIMAN	Т	T HI IMAN						T HUMAN	EST HUMAN	Т	T			
Top Hit Acession No.	K474R0n	3	T	1.0E-55 N77281 1	Š	0823120 BE077408 4	003.4.4	10077424	Ţ	4 0E-56 AF141340 1		1N 8777064	4507728 NT	003528.1	8924029	6912743 NT	TNI2897INT		3.0E-56 AA325828.1	Γ	393512.1	7657042	6326	5902085 NT	6912593 NT	6912593 NT	l		2.0E-56 BE064386.1	Γ	Γ	2	
Most Similar (Top) Hit BLAST E Value	1.0E-65	1 0F-65	1 0F-55	1.0E-55	1 0F 55	1 0F-55 RE	7 0F-58 H	6.0E-56 AV	4 0F-58	4 0F-56	4 OF 68	4 00 50	4.05-30	4.0E-56 AFC	3.0E-58	3.0E-56	3.0E-56	3.0E-56/	3.0E-56/	3.0E-56 AF0	3.0E-56 BE:	3.0E-56	3.0E-56 AL1	3.0E-56	3.0E-56	3.0E-56	2.0E-56 A	2.0E-56 B	2.0E-56 B	2.0E-56 M26061.1	2.0E-56 M26061.1	2.0E-56 A	
Expression Signal	1.83	3.86	1.04	101	108	22	3 83	183	44.14	44.14	67	2.7		3.56	3.78	3.22	1.19	1.47	1.47	2.03	1.28	1.01	4.75	2.24	0.72	0.83	1.87	1.67	1.67	0.94	0.94	1.04	
ORF SEQ ID NO:	13423	13882	14152		14913	l		11721	10092	10093	12709	12710		10560	11368	11800	12184	13078	13079		13827	14246	14279	14414	14898	14898	_	10765	10766	12416	12417	12955	
SEQ ID	8397	8884	9165	9580	9835	9866	7619	6849	6108	5108	7596	7596		5557	6320	6722	7070	8069	6908	8749	8820	9256	8292	9430	848	8918	6655	7729	7729	7296	7296	7838	
Probe SEQ ID NO:	3389	3883	4170	4592	4958	5015	2659	1653	28	28	2836	2636		2740	1322	1727	2089	3052	3052	3745	3818	4263	4300	4440	494 1	\$ 84	220	723	723	2321	2321	2919	

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	Top Hit Descriptor	Homo sablens gare for ectivin recentor has IIR complete ade	AV703184 ADB Homo sepiene cDNA clans ADBCEGO E	Macaca fascicularis protein tyrosine phosphatasa (PRI -1) mRNA commiste ode	254509.r1 Sogres overy tumor NbHOT Homo sepiens cDNA done IMAGE:726137 6' similar to gb:M94654	INTERNATION ENTENNESS BINDING FACIOR (HUMAN);	hazaciixi NCI COAP GCB Home seniens ADNA Alexanika CE 2014 24 201	QV0-0T0033-070300-162-h03 OT0033 Home seniens c-DNA	Homo saplens EphA4 (EPHA4) mRNA	Homo sepiens EphA4 (EPHA4) mRNA	QV4-ST0234-181199-037-f05 ST0234 Homo sepiens cDNA	x05d10.x1 NCI_CGAP_Bm53 Homo septens cDNA clone IMAGE:2759261 3' stmilar to go:U05876 INTERFERON-GAMMA RECEPTOR BETA CHAIN PRECI IBSOR ALI IMAAN	2051b12.11 Sogres testis NHT Hamp sapiens cDNA clane IMAGE-757151 5	Homo saplens EphA4 (EPHA4) mRNA	Homo sapiens EphA4 (EPHA4) mRNA	60094440F1 NIH_MGC_17 Homo septens cDNA clane IMAGE:2980884 5	Homo sapiens ang GDS-ASSOCIATED PROTEIN (SMAP), mRNA	Homo saplens sing GDS-ASSOCIATED PROTEIN (SMAP), mRNA	Homo sapiens NME7 (NME7), mRNA	Homo sapiens NME7 (NME7), mRNA	Homo sapiens Kruppel-like factor 8 (KLF8), mRNA	Homo sapiens phosphatidylinositol 4-kinase 230 (p4K230) mRNA, complete ods	Homo sapiens phosphatidylinositol 4-kinase 230 (p44K230) mRNA, complete cds	Homo sapiens FRA3B common fragile region, diadenosine triphosphate hydrolase (FHIT) gene, exon 5	Homo sapiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes, complete cds)	Homo saplens ubiquitin protein ligase E3A (fruman papilloma virus E6-associated protein, Angelman syndrome) (UBE3A) mRNA	nc13f07.s1 NCI_CGAP_Pr1 Homo sapiens cDNA clone IMAGE:1008037 similar to SW:RS10_HUMAN P46783 40S RIBOSOMAL PROTEIN S10 ;	EST54770 Hippocampus II Homo sapiens cDNA 5' end
	Top Hit Detabase Source	Z	EST HUMAN	۲	TOUR TOUR	EST HIMAN	EST HUMAN	Τ		1	EST_HUMAN	EST HUMAN	Ī			EST_HUMAN							뉟	_F	<b>−</b> ₽		EST_HUMAN	П
	Top Hit Acessian No.	AB008681.1	Γ	Γ		T		T	4758279 NT	4758279 NT	AW816405.1	AW 264599.1	Γ	4758279 NT	4758279 NT	3E299916.1	7657592 NT	7657592 NT	7242158 NT	7242158 NT	6005979	F012872.1	F012872.1	AF020503.1	AB026898.1	4507788 NT		3.0E-57 AA348335.1 E
	Most Similar (Top) Hit BLAST E Value	2.0E-56	2.0E-58	1.0E-56	4 OF 58	1.0E-56	1.0E-58	9.0E-57	9.0E-57	9.0E-57	8.0E-57	8.0E-57	_	8.0E-57	8.0E-57	8.0E-57	7.0E-57	7.0E-57	7.0E-57	7.0E-57	7.0E-57	7.0E-57 A	7.0E-57 A	7.0E-67	4.0E-57	3.0E-57	3.0E-57	3.0E-57
	Expression Signal	1.89	1.29	4.42	9 18	2.28	2.28	1.82	0.97	76.0	2.81	7.02	1.69	1.37	1.37	0.81	76.0	0.97	1.16	1.16	0.74	2.1	2.1	1.78	242	1.62	153.33	1.31
	ORF SEQ ID NO:		13491		11641)	L	13602		14063		10365	10832	11864	13333	13334	14825	12642	12843	13214	13215	13235	13793	13794		13677	10847		12421
	SEQ ID NO:		8464		6486			5647	9075	9075	5352	6891	6772	8308	8308	9848	7525	7626	8193	8183	8213	8789	8789	8300	8673	5816	6308	7301
Ī	Probe SEQ ID NO:	3242	3456	965	1489	3590	8	620	4081	4081	8	873	1780	297	3297	4869	2561	2661	3177	3177	3197	3788	3786	4308	3888	795	1311	2327

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	783910x1 NCI_CGAP_CLL1 Hamo saplens cDNA done IMAGE:3288443 3' similar to WP:Y47HpC.2 CE20283;	733b10x1 NCI_CGAP_CL.1 Homo sapiens cDNA clone IMAGE:3286443 3' similar to WP:Y47H9C.2 CE20263 :	Homo sapiens calline tsA2018 chloride ion criment inchiner protein I/On) cann complete ad-	RC3-CT0254-110300-027-d10 CT0264 Hamp septens CDNA	Homo sapiens SNARE protein kinase SNAK mRNA, complete ods	Homo sapiens SNARE protein kinase SNAK mRNA, complete cds	MR0-HT0559-010400-009-h10 HT0559 Homo sapiens cDNA	Homo seplens chromosome 21 segment HS21C004	ye98h01.r1 Sogres fetal liver spicen 1NFLS Homo sapiens cDNA clone IMAGE:125800 Fr	ye98h01.r1 Soares fetal liver spleen 1NFLS Homo seplens CDNA clone IMAGE-195ano R	MR0-BT0551-060300-103-b03 BT0551 Home saciens cDNA	2940c08.r1 Sogres retina N2b4HR Homo sapiens cDNA clone IMAGE:381450 R	2640c06.r1 Soares retina N2b4HR Homo sapiens cDNA clone IMAGE:361450 5	Homo sapiens chromosome 21 segment HS21C083	UI-HF-BNO-akt-g-07-0-UI.r1 NIH MGC 50 Homo sanians cDNA close IMAGE-3078348 F	601445948F1 NIH_MGC_65 Hamo saplens cDNA clane IMAGE:3850211 5	tr34b07.x1 NCI_CGAP_Ov23 Home sapiens dDNA clone IMAGE:2220181 3' similar to TR:016476 016476 UNNAMED HERV-H PROTEIN;	t34b07x1 NCI_CGAP_Ov23 Homo saplens cDNA clone IMAGE:2220181 3' similar to TR:015475 015475 UNNAMED HERV-H PROTEIN:	Homo saplens putative protein O-mannosytransferase (POMT2), mRNA	Homo sapiens putative protein O-mannosyltransferase (POMT2), mRNA	Homo saplens DHHC1 protein (LOC51304), mRNA	601309465F1 NIH_MGC_44 Home sapiens cDNA clone IMAGE:3831000 67	AU130689 NT2RP3 Homo sepiens cDNA clone NT2RP3001263 5	TCAAP1E1219 Pediatric acute myelogencus leukamia cell (FAB M1) Baylor-HGSC project≂TCAA Homo sapiens cDNA clone TCAAP1219	TCAAP1E1219 Pediatric acuts myelogenous leukemia cell (FAB M1) Baylor-HGSC project≃TCAA Homo saplens cDNA clone TCAAP1219	Homo sapiens synaptojanin 1 (SYNJ1), mRNA	RC4-NT0057-160600-016-b05 NT0057 Homo septens cDNA	CM3-UM0043-240300-127-e07 UM0043 Homo seplens cDNA
Top Hit Database Source	EST_HUMAN	EST HUMAN	Į.	EST_HUMAN	Z	Ę	EST HUMAN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	N-	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	5	17	トラ	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	1.	EST_HUMAN	EST_HUMAN
Top Hit Acessian No.	676622.1	676622.1		1853964.1	Γ			2			Г	Γ	2.0E-57 AA018299.1		1503208.1	868715.1	98376.1	98376.1	11434921	11434921 NIT	7706132 NT			6.0E-58 BE242150.1	6.0E-58 BE242150.1	7334		5.0E-58 AW797948.1
Most Similar (Top) Hit BLAST E Value	3.05-57 86	3.0E-57 BE	3.0E-57 AF	3.0E-57	2.0E-57	2.0E-57	2.0E-57	2.0E-57	2.0E-57	2.0E-57	2.0E-57	2.0E-57	2.0E-57	2.0E-57	1.0E-67 AW	8.0E-58 BE	8.0E-58 AI7	8.0E-58 AI7	8.0E-58	8.0E-58	8.0E-58	6.0E-58	8.0E-58	6.0E-58	6.0E-58	6.0E-58	5.0E-58	5.0E-58
Expression Signal	1.45	1.45	1.73	31.12	1.65	1.65	1.04	3.62	0.68	0.68	0.83	1.05	1.05	8.09	1.48	1.9	4.07	4.07	1.74	1.74	2.79	96.0	10.37	0.94	0.94	3.24	6.5	3.64
ORF SEQ ID NO:	12703	12704	13506		11526	11527	12432		13501	13502	13837	14044	14045	14349	12266		10679	10680	11902	11903		12292	12410	12870	12871	10367	·10740	11213
Exam SEQ ID NO:	7691	7591	8480		6468	6468	7311			╻┃	8830	9057	8057	9370	7149	5615	5674	5674	6810		١		7289	7851	7851	5355	5724	6178
Probe SEQ ID NO:	2631	2631	3482	3614	1471	1471	2337	3354	3474	3474	3828	4083	4063	4379	2170	\$	. 848	646	1820	1820	2806	2192	2314	2831	2831	298	200	1175

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	Top Hit Descriptor	CM3-UM0043-240300-127-e07 UM0043 Homo sapiens cDNA	CM3-UM0043-240300-127-e07 UM0043 Hamo sapiens cDNA	CM3-UM0043-240300-127-e07 UM0043 Homo expiens cDNA	or98e07.s1 NCI_CGAP_Lu5 Hamo sepiens cDNA clane IMAGE:1903908 3'	ts89e07.x1 NCI_CGAP_GC9 Hamo sapiens cDNA clone IMAGE:2238468 3' strutar to SW:PRO2_ACACA P19984 PROFILIN II;	Homo sapiens ATP synthese, H+ transporting, mitochandrial F1 complex, O subunit (digomycin sensitivity conferring protein) (ATP60) mRNA	Homo sapiens interleukin 10 receptor, beta (IL10RB), mRNA	Homo sapiens coagulation factor IX (plasma thromboplastic component, Christmas disease, hemophilia B) (F9) mRNA	Human beta-prime-adaptin (BAM/22) gene, exon 3	Homo sapiens EGF-like repeats and discoldin I-like domains 3 (EDIL3), mRNA	yg10e02.r1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMAGE:31693 5	Homo sapiens peptide YY (PYY) mRNA	602185789F1 NIH_MGC_45 Homo sepiens cDNA clone IMAGE:4309843 5	602185789F1 NIH_MGC_45 Homo sapiens cDNA clone IMAGE:4309943 5'	Homo saplens 5-aminolevulinate synthase 2 (ALAS2) gene, complete cds	eimiler to gb:X66391 60S r TAX responsive element binding	Human complement component C5 mRNA, 3'end	Homo sapiens NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 9 (22kD, B22) (NDUFB9), mRNA	EST369252 MAGE resequences, MAGD Homo septens cDNA	EST369252 MAGE resequences, MAGD Homo sepiens oDNA	Homo sapiens partial AF-4 gene, exons 2 to 7 and Alu repeat elements	hy10f08.x1 NCI_CGAP_GC8 Home saplens cDNA clone IMAGE:3196835 3'	Homo saplens uncharacterized bone marrow protein BM038 mRNA, complete cds	Homo sapiens sterol regulatory element binding transcription factor 2 (SREBF2) mRNA	oz43h01.x1 Soares_NhHMPu_S1 Homo sepiens cDNA clone IMAGE:16781293'	Homo sapiens TATA box binding protein (TBP) mRNA	601458531F1 NIH_MGC_66 Homo sepiens cDNA clone IMAGE:3882086 6
	Top Hit Database · Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	 	F7		Σ	Ę	EST_HUMAN	Þ	EST_HUMAN	EST_HUMAN	NT	EST HUMAN	L	— <b>Է</b>	EST_HUMAN	EST_HUMAN	NT	EST_HUMAN	NT	크	EST_HUMAN	5	EST HUMAN
	Top Hit Acession	W797948.1	W797948.1	N797948.1	1988183.1	636745.1	4502302 NT	4504634 NT	4503648 NT	36251.1	5031680 NT	R17879.1	1	BF569848.1	3.0E-58 BF569848.1		2.0E-58 BE208532.1		6274549 NT	W957182.1		1.0E-58 AJ238093.1		F217514.1	4759169 NT	1141063.1	4507378	F035327.1
	Most Similar (Top) Hit BLAST E Veitue	5.0E-58 A	5.0E-58 A	6.0E-58	5.0E-58 A	5.0E-58 A	4.05-58	4.0E-58	4.0E-88	4.0E-68 U	4.0E-58	3.0E-58 R	3.0E-58	3.0E-58 B	3.0E-58	2.0E-58	2.0E-58	1.0E-58 M	1.0E-58	1.0E-58 A	1.0E-58		1.0E-58	1.0E-58 A	1.0E-58	1.0E-58 A		6.0E-59 B
	Expression Signal	3.64	2.81	2.81	9.15	96:0	8.26	1.73	1.06	2.19	1.09	1.23	1.98	3.33	3.33	7.87	27.01	0.84	9.81	1.04	1.04	3.35	1.46	96'0	1.98	5.66	69.17	2.96
	ORF SEQ ID NO:	11214	11213	11214	13283	14114	10435		11494	12640	13666		11412	13138	13139	10977		10762	11089	11352	11353	11419			12801	14778	12264	
	Exam SEQ ID NO:		6178	6178	8262	9131	5421		6437	7524	8661		සිසි	8121		5943	6289	5736	0909			6370			7687	9795		7714
	Probe SEQ ID NO:	1175	1178	1176	3249	4136	372	788	1440	2559	3655	333	1366	3105	3105	976	1271	712	1051	1307	1307	1373	1624	2582	2730	4811	2168	177

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	81931-05-x1 Schneider fetal brein 00004 Home sentens cDNA cirms IMAGE-2783868 3' stenilos se	TR:075786 075786 GANGLIOSIDE-INDUCED DIFFERENTIATION ASSOCIATED PROTEIN 1	8493h05.x1 Schneider fetal brain 00004 Homo sepiens cDNA clone IMAGE:2783865 3' similar to TR-075788 075788 0ANGI IOSIDE NEU IOSI DIESEDENTIA ATOM OCO.	W48c11x1 Spares NF T GRC S1 Home seminar CDNA Account MA CERCERCAS		Human mRNA for KIAA0184 gene, partial cds	Homo sapiens phosphatidylinosibil-4-phosphate 5-kinase, type II, beta (PIPGK2B) mRNA, and translated products	Homo seplens phosphatidylinositol 4-phosphate 5-kinase, type II, beta (PIP5K2B) mRNA, and translated products	W832612 XT NCL CRAP GCS Homo semiens of NA Alexa MA CE SACOSOS of	EST377682 MAGE resentences. MAGI Home contens milks	Homo sabiens KIAA0880 para product (KIAA0880) DNA	Homo saplens plasminoden activator, fissue (PLATs) mRNA	Homo saplens plasminoden activator, tissue (PLATs) mRNA	Homo sapiens mRNA for KIAA1112 protein, partial cds	Homo sapiens mRNA for KIAA1112 protein, partial cds	Homo saplens NF1-2 pseudogene, excn 17	Homo sapiens A kinase (PRKA) anchor protein 1 (AKAP1), mRNA	Homo sapiens A kinase (PRKA) anchor protein 1 (AKAP1), mRNA	Homo sapiens zona pellucida giycoprotein 2 (sperm resentor) (7P2) mRNA	Homo sapiens chromosome 21 segment HS21C084	Homo seplens protein tyrosine phosphatese, receptor type, T (PTPRT), mRNA	Human prohormone converting enzyme (NEC2) gene, exon 2	601176757F1 NIH_MGC_17 Homo septens cDNA clone IMAGE:3531927 5'	0956h11.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1309029 3' similer to TR:Q13637 O13537 MER37 TRANSPOSABLE ELEMENT. COMPLETE CONSENSUS SEOLIENCE	EST389849 MAGE resequences, MAGO Homo saplens cDNA	Homo saplens small nuclear ribonucleoprotein D3 polypeptide (18kD) (SNRPD3) mRNA	Homo saplens differentiation-related gene 1 (nickel-specific induction protein) (RTP) mRNA			Homo sapiens MHC class 1 region
Top Hit Datzbase Source		EST_HUMAN	EST HUMAN	Т	Т				T HUMAN	Т				LN		¥				TN L		NT	EST_HUMAN (	EST HUMAN	EST_HUMAN					
Top Hit Acession No.		AW157281.1	5.0E-59 AW157281.1	-			4505818 NT	4505818	980847.1	W965524.1	2247	4505860 NT	4505860 NT			3.0E-59 AF232299.1	4502014 NIT	4502014 NT	4508044 NT	163284.2	7427522 NT		1.0E-59 BE296411.1	1.0E-59 AA748468.1	8.0E-60 AW977845.1	4759159 NT	5174656 NT	4656	7.0E-60 AF055068.1	
Most Similar (Top) Hit BLAST E		5.0E-59 A	5.0E-59	5.0E-59	5.0E-59 X83497.1	4.0E-59	4.0E-59	4.0E-59	4.0E-59 A	3.0E-59/	3.0E-59	3.0E-59	3.0E-59	3.0E-59	3.0E-59	3.0E-59 /	3.0E-59	3.0E-59	3.0E-59	3.0E-59 A	3.0E-59	3.0E-59 M95961.1	1.0E-59	1.0E-59	8.0E-60	8.0E-60	8.0E-80	8.0E-60	7.0E-60 /	7.0E-60 A
Expression Signal		9.16	9.16	6.86	9.33	2.84	0.67	0.67	0.98	4.74	4.43	8.3	8.3	5.59	5.59	. 0.98	3.67	3.67	1.33	1.09	1.64	0.92	37.68	2.32	2.17	8.32	1.59	1.59	33.65	109.11
ORF SEQ ID NO:		11788	11787	13081	14498	10837	11258	11259	14964		10295	11743	11744	12162	12163	12769	13085	13086	13738	14523	14662				10803	11497	12201	12202	10794	10794
Exan SEQ ID NO:		6710	6710	1408	9513	5806	6217	6217	8888	2080	5287	8999	8999	7053	7053	77.0	8073	8073	8739	9236	9678	88	5228	7509	5778	6440	7087	7087	6768	5768
Probe SEQ ID NO:		1715	1715	3054	4523	785	1218	1218	5017	10	225	1672	1672	2071	2871	2697	3056	3056	3735	4547	4693	4884	162	2544	754	1443	2107	2107	745	748

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	801300838F1 NIH MGC 21 Home savious along the Classical III A	60/300938F1 NIH MGC 21 Home series about the MACE 3033480 5	Homo sablens PRO2014 mRNA complete Ale	Infeth09.st NCI CGAP Lart Home sapiens child close Mile CE-messes 2:	Home entire ed to contact (St Postate) and	AL130689 NT38P3 Home earlies - CANA - Intropose cas; nuclear gane for mitochandrial product	Homo sapiens T-cell humbroms triverion and motochasts of (TIANA). Days	Homo satients profess between a remindence of the property of the profession of the	Homo sapiens chromosome 21 segment HS210779	Homo saniens amwied hats (AA) macrinese executes (see the second of the	Hamp sapiens 959 kh contra helysees AMT 4 and CPB1	Homo sepiens T-cell banchome truesion and make less 4 (TIA14). This	Homo sapiens hypothetical protein El 14008 /El 144000 - Data	QV3-HT0513-060400-147-d01 HT0513 Home services CDNA	QV3-HT0513-060400-147-d01 HT0513 Home sealens CONA	W53d11.s1 Soares fetal liver spiesn 1NFLS Home sapiens CDNA clone IMAGE:248463 3' similar to obi 25444 60S RIBOSOMAL PROTEIN 1 4 to ALI IMAAN.	W03(11.r1 Soares melanocula 2NHHM Homo septems CNN close IMAGE:577450 2:	Homo sablens chromosome 21 segment HS21Ch03	Homo saplens origin recognition complex submit 2 (west homology Like (ADC) 1 - DNA	Human polymorphic trinucleotide repeat in X-linked retinitis plamentres (RPA) gene market	Homo sapiens zona pellucida glycoprotein 34 (sperm receptor) (ZP3A), mRNA	xn11b09.yi NCI_CGAP_LI5 Homo septens cDNA clone IMAGE:2693369 5' similar to contains element	601273513F1 NIH MGC 20 Homo serviens -DNA - Jone 11/4 OE-2841867 F1	Homo sablens KIAA0808 days profiled (KIAA0808) mRNA	Homo sepiens TRAF family member-associated NFKR activator (TANK) mbna	Homo saplens TRAF family member-associated NFKB activator (TANK) mRNA	UI-H-BW0-git-b-08-0-UI.s1 NCI_CGAP_Sub6 Homo sapiens cDNA clone IMAGE:2733871 3	UI-H-BW0-git-b-08-0-UI.s1 NCI_CGAP_Sub6 Hamo saplens cDNA clane IMAGE:2732871 9'	oc66h11.s1 NCt_CGAP_GCB1 Homo saplens cDNA clone IMAGE:1354725 3' similar to SW:POL_MLVRK P31795 POL POLYPROTEIN ;
Top Hit Databerse Source	EST HUMAN	EST HUMAN	N	EST HUMAN	· - 5	EST HIMAN	L		Z	 	ĬN		=	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	NT	Ļ	NT	나	EST HIMAN	EST HUMAN	-	7	5	EST_HUMAN	EST_HUMAN	EST_HUMAN
Top Hit Acessian No.	BE408310.1	Γ	Γ	Γ		T	2750	4506008 NT	4L163279.2	4502168 NT	AJ229041.1	7500		168410.1	168410.1	33039.1	39397.1	2	5453829 NT	J32657.1	F005983 NT	1.0E-81 AW827281.1	T	7662319	4759249 NT	4759249 NT	1.0E-61 AW298181.1	1.0E-61 AW 298181.1	8.0E-62 AA830420.1
Most Similar (Top) Hit BLAST E Value	8.0E-81	6.0E-81	6.0E-61	8.0E-61	8.0E-81	6.0E-61	5.0E-61	6.0E-61	5.0E-61	5.0E-61	5.0E-81	5.0E-61	2.0E-61	2.0E-61	2.0E-61 BE	2.0E-61 N	2.0E-61	1.0E-61	1.0E-81	1.0E-61	1.0E-61	1.05-61	1.0E-61 B	1.0E-61	1.0E-61	1.0E-61	1.0E-61	1.0E-61 /	8.0E-62
Expression Signal	4.42	2.26	12.89	2.82	0.95	13.67	0.88	3.51	2.15	1.7	1.66	1.18	1.33	3.21	3.21	1.75	1.38	0.75	1.16	0.96	4.18	1.82	2.42	0.73	0.75	0.75	9.13	9.13	0.92
ORF SEQ ID NO:	10331		11345	11668	12/58	13270	10420	11706	12997	13162		10420	10538	11231	11232	11692			10814		11804	12227	12807	13328	14294	14295	14676	14677	14388
Exan SEQ ID NO:	5322	5823	6233	8605	7050	8249	5408	6636	7983	8141	8875	5408	6531	6184	6194	6623	7533	5470	5785	6727	6811	7114	7782	8302	8309	8309	8693	8693	9410
Probe SEQ ID NO:	263	802	1301	1609	2068	3234	358	1639	2865	3125	3874	4843	495	1193	1193	1626	2570	432	764	1732	1821	2134	2761	3291	4317	4317	4708	4708	4420
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Single Exon Probes Expressed in HBL100 Cells

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Single Excite Forces Expressed in right come	Top Hit Descriptor	AV714334 DCB Hama sepiens cDNA dane DCBAMA08 5	NUCLEOLAR TRANSCRIPTION FACTOR 1 (UPSTREAM BINDING FACTOR 1) (UBF-1) (AUTOANTIGEN NOR-60)	Human zinc finger protein ZNF131 mRNA, partiel cds	Homo sapiens CGI-56 protein (CGI-66), mRNA	w.51e07.x1 NCI_CGAP_Lu28 Homo sapiens cDNA clone IMAGE:2547204 3' similar to SW:GG95_HUMAN Q08379 GOLGIN-05, ;contains element MER22 repositive element;	Homo sapiens Xq pseudosutosomal region; segment 1/2	Homo sapiens Xq pseudosutosomal region; segment 1/2	Human xanthine dehydrogenase/oddase mRNA, complete ods	Human xanthine dehydrogenase/oxidase mRNA, complete cds	Homo sepiens ryanodine receptor 3 (RYR3) mRNA	zw78e09.s1 Soares_testis_NHT Homo sepiens cDNA clone iMAGE:782344 S' similar to SW:NRDC_RAT P47245 NARDITYSIN;	IRC5-IN1089-100500-021-H03 NN1089 Homo sepiens cDNA	Bu71403.y1 Schneider fetal brein 00004 Homo sepiens CDNA clone IMAGE:2781701 6' amitar to go:n437104   ATP SYNTHASE COUPLING FACTOR 6, MITOCHONDRIAL PRECURSOR (HUMAN);	au74d03.y1 Schneider fetal brain 00004 Homo saptens cDNA clone IMAGE:2781701 6' similar to gb:M37104	ALT STATIS OF Schrades feels from Anna Banishs CDNA close IMAGE:2781701 5 similar to ab:M37104 Fe	ATP SYNTHASE COUPLING FACTOR 6, MITOCHONDRIAL PRECURSOR (HUMAN);	au/1403.y1 Schneider fetal brain 00004 Homo septens CINA clone IMACE:2/61/01 5 SIMMET to grams / 104 ATP SYNTHASE COUPLING FACTOR 6, MITOCHONDRIAL PRECURSOR (HUMAN);	wf12b08.x1 Soares_NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE:2350359 3' similar to qb:X57139_me1 HISTONE H2B.2 (HUMAN);	w112b08.x1 Sogres_NFI_T_GBC_S1 Homo sapiens cONA clone IMAGE:2350359 3' similar to	gb:X57138_ma1 HISTONE H2B.2 (HUMAN);	Homo sapiens keratin 18 (KRT18) mRNA	Homo saplens enhancer of zaste (Drosophila) homotog z (E.Z.H.z) myrink	Homo sapiens neurofibromin 2 (bilateral acoustic neuroma) (NFZ) mKNA	Homo sapiens mRNA for KIAA1476 protein, partial cds	l cds		Homo sapiens chromosome zi segment nozi coor
Jeseph Ling	Top Hit Database Source	EST HUMAN	SWISSPROT	NT	11	EST_HUMAN	NT	NT	LN	NT	17	EST_HUMAN	EST_HUMAN	EST HUMAN		HOWAN	EST_HUMAN	EST HUMAN	EST HUMAN		EST HUMAN	NT	LN	NT	NT.	NT	NT	<u>F</u>
סווולווס	Top Hit Acessian No.	714334.1	7480		11418255 NT	6.0E-62 Al950528.1			5.0E-62 U39487.1	5.0E-62 U39487.1	4506758 NT	5.0E-62 AA431083.1				4.0E-62 AW1614/9.1	4.0E-62 AW161479.1	4.0E-62 AW161479.1			41827800.1	4557887	4758323 NT	4557794 NT	AB040909.1	AB040909.1	X52858.1	AL163284.2
	Most Similar (Top) Hit BLAST E	7.0E-62 A\	7.0E-62 P	6.0E-62 U09410.1	6.0E-62	6.0E-62	6.0E-62	6.0E-62	5.0E-62	5.0E-62	5.0E-62	5.0E-62	5.0E-82	4.05.63		4.0E-82	4.0E-62	4.0E-62	4.0F-62		4.0E-62	4.0E-82	4.0E-62					2.0E-62
	Expression Signal	1.09	860	1.66	4.7	3.65	3.43	3.43	96'0	0.98	2.46	1.82	1.12	4 H	5	5.18	3.63	8	577		4.43	7.88	0.97					2.89
	ORF SEQ ID NO:	11126				10470						<u> </u>		10007		10888	10887		<u> </u>		12483		14967		ľ			11251
	Esan SEQ ID NO:	6097	8435	7849	8311	5450	7315	7315	7483	7483	8344	9194	9417	0202		5850	6850	ABEO	7360	5	7360		L				1	
	Probe SEQ ID ( NO:	1000	3427	2930	3300	413	2341	234	2515	2515	3334	4201	4427	8	3	88	83	8	Š	ROCZ	2380	3315	5023	2	2072	2972	3615	1211

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	Top Hit Descriptor	Homo sanions Intersectin 3 /SH3D4B) = DNA	Homo septems ADP/ATP center protein (ANT-3) mans	#70e11.r1 Soares_NiHMPu_S1 Home septens CONA clone IMAGE:1047404 6' stmiler to WP:K01H12.1	0/67-588-404 -4 Fee /	Disk 200001104_11 500 (synon)m: rinkd2) Homo sapiens cDNA clone DKFZp668F104 6'	Homo saplens hypothetical protein FL/20212 (FL/20212) mRNA	206b08.r1 Soeres_pregnant_uterus_NbHPU Homo saptens cDNA clone IMAGE:491611 6 similar to SW:CS81 BOVIN P10897 CYTOCHEOME BRAS	OV4-ST0234-181109-037-815 ST0234 Domp contons - PATA	C18159 Human placenta cDNA (TELithware) Homo emission colla state process	Homo sablens mRNA for KIA AARA models and the septents of the CEN-558CTO 6	Homo saplens mRNA for KIAA0350 profein partial cde	2831408-r1 Soures reithe N2b4HR Home sapiens CDNA done IMAGE:360561 6" similar to SW:UN13_CAEEL P27715 PHORBOL ESTERDACH RI YERRI JRINDING BEATHERN 1902 223	Hymn serviews management at 11.000 to 11.000 t	Homo earloss (1.2 Indicate in the Contract of	Gallus nating Darb? protein (Darbs) mDNA completed	Gallus callus Dach? motein (Dach?) mRNA complete ada	Homo saplans chromosome 21 semment HS94 Chea	wm55g11.x1 NCI CGAP Ut2 Home sablens cDNA refere IMA GE-2490008 9	Homo saplens chromosome 21 segment HS21C078	Homo sapiens mRNA for KIAA0707 protein, partial cds	Homo sapiens mRNA for KIAA0707 protein, partial cds	Homo sapiens mRNA for KIAA0717 protein, partial cds	Human Met-IRNA-I gene 1	Homo sapiens zinc finger protein 144 (Met-18) (ZNF144), mRNA	Human DNA topotsomerase I mRNA, pertial cds	Homo sapiens eyes absent (Drosophila) homolog 2 (EYA2), mRNA	Homo sapiens giutamata-cysteine ligase (gamma-giutamylcysteine synthetase), catalytic (72.8kD) (GLCLC) mRNA	Homo saplens Down syndrome candidate region 1 (DSCR1), mRNA	Homo sapiens RHCE mRNA for Rh blood CE group antigen polypeptide, complete cds
89701 1 10%	Top Hit Database Source	L	Į.	EQT LINKAN	EST HIMAN		TN	EST HUMAN	EST HUMAN	EST HUMAN	LN	¥	EST_HUMAN		5	Z	날	Ŀ	EST HUMAN	L.	Ę	NT	NT	NT		Į.				LN L
	Top Hit Acession No.	1.0E-62 AF248540.1			AI 039044 1	1.0E-62 AB040911.1	8		Г	Γ		9.0E-63 AB002348.2		4557734 NT	5031810 NT	98349.1	98349.1	63268.2	72137.1			1.	.1		TN 5963009	804.1	4885226 NT	4557624 NT	7857042 NT	30388.1
	Most Similar (Top) Hit BLAST E Value	1.0E-62	1.0E-82	1.05-82	1.0E-82	1.0E-62	1.0E-62	1.0E-62	9.0E-83/	9.0E-83	9.0E-63	9.0E-83 /	9.0E-63 AA	8.0E-63	8.0E-63	8.0E-63 AF	8.0E-63 AF1	8.0E-63 AL	7.0E-63	4.0E-63 AL1	4.0E-63	4.0E-63 A	3.0E-63 AB018260	3.0E-63	3.0E-63	2.0E-63 U07	2.0E-63	2.0E-63	2.0E-83	2.0E-63 AB0
	Expression Signal	1.58	15.3	1.92	1.18	2.49	1.63	0.98	1.82	1.09	9.26	9.28	4.71	2.39	5.14	4.89	4.89	3.27	1.84	0.7	2.01	2.01	2.67	1.34	10.16	2.47	1.85	1.36	4.72	1.52
	ORF SEQ ID NO:	11069	11587	11840	12882		14369	14911	10395		13918	13919	14995	12376	12409	13408	13409	14122		13282	. 13722	13723	11975	12774	11262	10267	10275		10872	11591
	Exan SEQ ID NO:	6037	6510	6755	7864	8348	9386	9933	6388	7260	8928	8928	10026	7258	7288	8387	8387	9138	5932	8260	8723	8723	883	7683	6219	5255	6262	5530	5837	6531
	Probe SEQ ID NO:	1027	1512	1763	2844	3338	4395	4956	338	2284	3928	3928	5056	2282	2313	3379	3379	4143	916	3247	3719	3719	1895	2706	2748	ᅙ	188	494	816	- E

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Single Exoli Flores Expressed III IDE IO Colis	Top Hit Descriptor	Homo sepiens RHCE mRNA for Rh blood CE group antigen polypeptide, complete ods	601301627F1 NIH_MGC_21 Hamo septens cDNA clane IMAGE:3638103 6	Homo sapiens amyloid beta (A4) precursor protein (protease nextr-II, Alzheimer disease) (APP), mRNA	Homo sapiens chromosome 3 subtelomeric region	Homo saplens polycystic kidney disease-associated protein (PKD1) gene, complete cos	Homo sapiens jun dimertzation protein gene, pertial cds; cfos gene, complete cds; end unknown gene	HSCZVD111 normalized infant brain cDNA Homo sepiens cDNA clone c-zvd11	HSCZVD111 normalized Infant brain cDNA Homo sapiens cDNA clone c-zvd11	601155232F1 NIH_MGC_21 Hamo sepiens cDNA clone IMAGE:3139038 6	601311455F1 NIH_MGC_44 Homo saplens cDNA clone IMAGE:3633204 5	Homo sapiens thimet oligopeptidase 1 (THOP1) mRNA	Homo sapiens thimet oligopeptidase 1 (THOP1) mRNA	wb51e07.x1 NCI_CGAP_GC8 Hamo septens cDNA clone IMAGE:2308220 3' similer to gb:M15182 BE LA- GLICURONIDASE PRECURSOR (HUMAN);	TESTOT A NO. COAD COR Dame contains a DNA riems INAGE: 2309220 3' similar to oth-M 5182 BETA-	WESTELL'S INCLUSION THE SEPTING SEPTIN	wv13e03.x1 NCL_CGAP_Brn23 Homo sepiens cDNA clone IMAGE:2528436 3	w/13603.x1 NCI_CGAP_Bm23 Homo sapiens cDNA clone IMAGE:2529436 3	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens mRNA for KIAA0903 protein, partial cds	Home sapiens phosphoglucomutase-related protein (PGMRP) gene, complete cds	Homo sepiens phosphogiucomutase-related protein (PGMRP) gane, complete cus	Human (3)mbt protein homolog mRNA, complete cas	Homo sapiens KIAA0618 gene product (KIAA0618), mKNA	Homo sapiens KIAA0618 gene product (KIAA0618), mRNA	Homo sapiens putative transcription factor CR53 (CR53) mRNA, partial cds	Homo sapiens mRNA for KIAA0903 protein, partial ods	C18895 Human placenta cDNA (TFujiwara) Homo saplens cDNA clone GEN-568E02 5	AV711714 DCA Homo seplens cDNA clone DCAAMC01 6	AV711714 DCA Homo sapiens cUNA cione DCAAMICUT o	BUUSAUS I SOBRES LES LES LES LES LES LES LES LES LES L
	Top Hit Database Source		EST. HUMAN	•	ı ı			EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN			NAMINAN		EST_HUMAN	EST_HUMAN	EST_HUMAN	_		1	ı	L	_	T	1	1	ĮN.	EST_HUMAN	ST HUMAN	EST HUMAN	EST HOMAN
3	<u> </u>	Ā	S	36 NT	INT	Ę	Ę	ES	ES	ES	ES	N 06	IN 06	ŭ	4	. Ш	ä	ĬŬ.	Z	IN	Z	¥	Z	Ŋ	7662205 NT	7662205 NT	호	Z	Ш	3	9	4
Sirgie	Top Hit Acessian No.	B030388.1	2.0E-63 BE410739.1	4502166 NT	₹109718.1	2.0E-63 L39891.1	2.0E-63 AF111167.2	1.0E-83 F08485.1	1.0E-63 F08485.1	8.0E-64 BE280798.1	7.0E-64 BE394321.1	4507490 NT	4507490 NT	A 1884 000 4		6.0E-64 Ai651992.1	AW026445.1	AW026445.1	14	AF231919.1	AB020710.1	L40933.1	L40933.1	U89358.1			AF017433.1	AB020710.1				AA609940.1
	Most Similar (Top) Hit BLAST E Value	2.0E-63	2.0E-63	2.0E-83	2.0E-63	2.0E-63	2.05-63	1.0E-63	1.0E-63	8.0E-64	7.0E-64	7.0E-84	7.0E-64	4 10 0	0.0E-04	6.0E-64	6.0E-64	6.0E-84	5.0E-64	5.0E-64	5.0E-64	5.0E-64	5.0E-64	5.0E-64	5.0E-64	5.0E-84			3.0E-64	3.0E-64		2.0E-84
	Expression Signal	1.52	3	1.58	22	1.4	1.23	3.33	3.33	9.14	6.0	3.25	3.25	, a	G.43	6.45	609	5.09	3.24	3.24	2.38		1.35	1.5	4.17							1.32
	ORF SEQ ID NO:	11592	11802	13114	13249		14887					14570	14571		20/5	11764		L			١							L	L			1111
	Exan SEQ ID NO:	6831	6724	808	8227	8812	9701	8209	9209	8039	8459	9581	9581	8	200	6880	ROBB	8086	5831	583	6316	6394	6394	6999	6447	8447	8854	8988	7115	8367	8367	6080
	Probe SEQ ID NO:	1633	1728	3083	3212	3809	4716	4216	4216	1029	3451	4583	4593		1684	1684	3040	8040	340	a Q	13/8	1397	1397	1673	2753	27.53	3853	808	24.85	3350	3359	1072

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Table 4
Single Exon Probes Expressed in HBL100 Cells

w87b01.x1 NCI\_CGAP\_Kid11 Homo sepiens cDNA clone IMAGE.2462281 3' sémilar to contains element Homo sapiens glutamic-oxaloacetic transaminase 2, mitochondrial (aspartate aminotransferase 2) (GOT2), nuclear gene encoding mitochondrial protein, mRNA Homo sapiens transcription factor IGHM enhancer 3, JM11 protein, JM4 protein, JM5 protein, T54 protein, qm48e01.x1 Soares\_placents\_8tu8weeks\_2NbHP8to9W Homo explens cDNA clone IMAGE:1891800 3\* qm46e01.x1 Soares\_placenta\_8to9weeks\_2NbHP8tc9W Homo seplans cDNA clone IMAGE:1891800 3\* JM10 protein, A4 differentiation-dependent protein, triple LIM domain protein 6, and synaptophysin genes, nj86d10,s1 NCI\_CGAP\_Pr11 Homo sapiens cDNA clone IMAGE:999379 similar to gb:K03002 60S au60cc1.x1 Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2519138 3' similar to gb:L21696\_cds1 PROTHYMOSIN ALPHA (HUMAN);contains element MSR1 repetitive element; DKFZp761G108\_11761 (synonym: hamy2) Homo sapiens cDNA clone DKFZp761G108 6 Homo septens fregile X mental retardation, autosomel homolog 1 (FXR1), mRNA Homo septens ribosomel protein L34 (RPL34) mRNA Homo sepiens ubiquitin specific protesse 13 (isopsptidase T-3) (USP13) mRNA Homo sepiens ubiquitin specific protesse 13 (isopsptidase T-3) (USP13) mRNA Homo sapiens hypothetical protein FLJ11028 (FLJ11028), mRNA Top Hit Descriptor formo septems el F4E-like cep-binding protein (4EHP) mRNA Homo sapiens KIAA0156 gene product (KIAA0168), mRNA Homo sapiens KIAA0156 gene product (KIAA0156), mRNA AV721898 HTB Homo sapiens cDNA clone HTBBZC06 5 H. sapiens DNA for endogenous retroviral like element Homo sapiens chromosome 21 segment HS21C048 Homo sapiens chromosome 21 segment HS21C046 Homo sapiens chromosome 21 unknown mRNA Homo seplens KE03 protein mRNA, partial cds Homo sapiens synaptojenin 1 (SYNJ1), mRNA H.sapiens DNA for endogenous retroviral like complete cds; and L-type calcium channel a> Homo sepiens TRIAD3 mRNA, partial cds Homo sepiens TRIAD3 mRNA, partial cds RIBOSOMAL PROTEIN L32 (HUMAN); L1 repetitive element EST HUMAN EST HUMAN EST\_HUMAN EST\_HUMAN EST HUMAN EST HUMAN **EST HUMAN** Top Hit Database Source Ę Ę 7881951 NT 4757701 NT 4504068 NT 늘 Z 4506636 NT 4507848 N 1.0E-64 AI828419.1 4507334 N 4826735 8922829 7661951 Top Hit Acession 4507 2.0E-64 AL163246.2 5.0E-65 AA550929.1 2.0E-64 AI927030.1 2.0E-64 AL163246.2 1.0E-64 AF231919.1 AL120419.1 .0E-64 AF196779.1 1.0E-64 AF228527.1 4.0E-65 AI266468.1 AV721898. 4.0E-65 AI266468.1 .0E-64 AF228527. 9.0E-65 X89211.1 9.0E-65 5.0E-65 5.0E-85 4.0E-65 1.0E-64 6.0E-65 6.0E-65 4.0E-85 2.0E-64 5.0E-65 (Top) Hit BLAST E 2.0E-84 6.88 18.94 1.92 51.83 0.81 48.55 1.07 5.42 2.92 1.58 1.09 1.09 0.98 <u>4</u> 1.94 8. 1.07 <u>9</u> 0.87 3.17 Expression Signal 11812 13814 12312 10856 11381 11382 1078<u>2</u> 1110<u>2</u> 11420 12544 12545 13097 13526 11079 11511 10325 13526 10781 12981 13221 ORF SEQ 1027 Ö Z Q SEQ ID 7424 8083 7961 6989 6333 5759 5759 6070 6452 6371 7428 5316 6333 8197 888 7191 **8**81 5257 851 ÿ 736 2454 2459 3432 2214 1039 1880 625 1335 1335 3181 1455 Probe SEQ ID 3503 3805 736 2459 256 3181 3067 1741 2942 2214 1374 ÿ

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	hu25e04.x1 NCI_CGAP_Me15 Homo septems cDNA clane IMAGE:3171102.3*	hu25604.xf NCI CGAP Mel15 Homo explans abiNA alone IMAGE:3171102 3"	RC2-BN0033-160200-013-603 BN0033 Homo saciens cDNA	Homo sepiens pre-B-cell colony-enhancing factor (PBEF) mRNA	Homo saplens pre-B-cell colony-enhancing factor (PBEF) mRNA	H. sapiens HZF9 mRNA for zinc finger protein	Homo sapiens immunoglobin superfamily, member 3 (IGSF3) mRNA, and translated products	ov23f03.s.1 Soares_bests_NHT Homo septens cDNA clone IMAGE:1638173 3' similar to contains element MSR1 repetitive element:	Homo sapiens mRNA for KIAA0235 protein, pertial cds	Homo septens laminin, beta 1 (LAMB1), mRNA	ov23f03.s1 Soarss_testis_NHT Homo sapiens cDNA done IMAGE:1638173 3' similar to contains element MSR1 repetitive element;	Homo eaplens rabs GTPesse activating probeh (GAP and centrasome essociated) (QAPCFNA) mRNA	602155062F1 NIH MGC 83 Homo sabians cDNA clone IMAGE:4296988 67	601763488F1 NIH_MGC_20 Hamo saplens cDNA clone IMAGE:4026501 67	Homo saplens putative Rab5 GDP/GTP exchange factor homologue (RABEXS), mRNA	Homo sapiens mRNA for KIAA1513 protein, partial cds	hz24a09.x1 NCI_CGAP_GC8 Homo sepiens cDNA clone IMAGE:3208888 3'	Homo saplens glypican 4 (GPC4) mRNA	Homo sapiens glypican 4 (GPC4) mRNA	wx08c09.x1 NCI_CGAP_Gas4 Homo seplens cDNA clone IMAGE-2543152 3'	wx09c09.x1 NCI_CGAP_Gas4 Homo sapiens cDNA clone IMAGE:2543152 3'	Novel human gene mapping to chomosome 22	Novel human gene mapping to chomosome 22	Homo sapiens 26S proteascome-associated pad1 homolog (POH1) mRNA	Homo saplens 26S proteasome-associated pad1 homolog (POH1) mRNA	Human transposon-like element, partial	Novel human gene mapping to chamosame X	wn57h07.x1 NCi_CGAP_Lu19 Homo saplens cDNA clone IMAGE:2449597.3' similar to WP:F15G9.4A CE18595;	wn57h07x1 NCI_CGAP_Lu19 Hamo sapiens cDNA clone IMAGE:2449597 3' similar to WP:F15G9.4A CE18595;
	Top Hit Database Source	EST HUMAN	EST HUMAN	EST HUMAN	Z	¥	NT	NT	EST HUMAN	NT	NT	EST_HUMAN	_ <u></u>	EST HUMAN	EST HUMAN	NT	. IN	EST_HUMAN	NT	NT	<b>EST_HUMAN</b>	EST_HUMAN	NT .	NT	NT .	TN	TN	NT	EST_HUMAN	EST_HUMAN
,	Top Hit Acessian Na.	BE221469.1	BE221469.1	AW883185.1	5031976	5031976 NT	X78932.1	4504626 NT	A1000692.1		4504950 NT	A1000692.1	0912385 NT	BF680294.1	Γ	7657495 NT	AB040946.1	BE466681.1	4504082 NT	4504082 NT	AW029340.1	AW029340.1		AL160311.1	5031980 NT	5031980 NT		AL137163.1	AI924653.1	AI924653.1
	Most Similar (Top) Hit BLAST E Value	4.0E-85	4.0E-65	4.0E-65	3.0E-65	3.0E-65	3.0E-65	3.0E-65	3.0E-85	3.0E-65	3.0E-85	3.0E-85	3.0E-85	2.0E-65	1.0E-65	1.0E-85	1.0E-65	1.0E-65	1.0E-65	1.0E-65	1.0E-65	1.0E-65	9.0E-66	9.0E-88	9.0E-68	9.0E-68	9.0E-66	9.0E-66	6.0E-88	6.0E-68
	Expression Signal	3.62	3.62	1.07	1.82	1.75	12.12	1.55	1.62	0.89	0.83	1.17	1.45	6.2	1.12	1.79	1.12	0.79	1.89	1.89	3.43	3.43	1.28	1.28	2.47	2.47	3.8	96'0	1.02	1.02
	ORF SEQ ID NO:	12368	12369	13855	10185	10185		11581	11871	12958	13241	13650	14485	13351		10572	12079	13327	13890	13891	14068	14067	10158	10157	11383	11384		14526	14216	14217
	Exan SEQ ID NO:		7251	8847	5174	5174	1	6524	8229	7940	8218	8644	9505	8330	5166	6959	6974	8301	8892	8892	9078	8078	5149	5149	6334	6334	6450	9541	9234	9234
ſ	Probe SEQ ID NO:	2275	2275	3845	96	26	1212	1527	1787	2921	3203	3638	4515	3320	88	534	1889	3290	3892	3892	4084	4084	70	70	1336	1336	1453	4553	4240	4240

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Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	wn57h07.x1 NCI_CGAP_Lu19 Homo septens cDNA ctons IMAGE:2449567 3' stmlier to WP:F15G9.4A CE18595 ;	RC4-BT0311-141169-011-NG BT0311 Homo septens cDNA	601681592F1 NIH MGC 9 Homo septens cDNA clone IMAGE:3961781 6	601681592F1 NIH_MGC_9 Home septens CUNA cione IMAGE:3801781 5	Mus musculus fragile X mental retardation syndrome 1 homolog (rimin), mixtva	RC1-NN0063-100500-022-e02 NN0063 Homo septems cUNA	H.sapiens DNA for endogenous retroviral like element	Homo saplens germ-line DNA upstream of Jkappa locus	Human endogenous retrovirus, comprete genome	Homo sapiens solute carrier family 25 (mitochondrial <b>carrier; attentive nucleon</b> de transioceux), mentioer o (SLC25A6), nuclear gene encoding mitochondrial protein, mRNA	Homo expiens solute carrier family 25 (mitochondrial carrier; aderane nucleotide translocator), member 5 (St. C25A5), nuclear gene encoding mitochondrial protein, mRNA	v27012.r.f Soares multiple sclerosis 2NbHMSP Homo sepiens cDNA clone IMAGE:284326 6' stmilar to	SW:H281_TIGCA P35068 HISTONE H2B.1/H2B.2. [2] PIR:B56612;	y27g12.rt Scares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA clone IMAGE:284328 6' similar to	SW:H2B1_TIGCA P35068 HISTONE H2B.1/H2B.2. [2] PIR:B56612;	y27g12.r1 Sogres_multiple_sclerosis_2NbHMSP Homo sapiens cuna cigne image:2003.c0 5 minum to SW:H2B1_TIGCA P35068 HISTONE H2B.1/H2B.2. [2] PIR:B56612;	Homo sapiens TGF(beta)-Induced transcription factor 2 (TGIF2), mRNA	Homo sapiens KIAA0649 gene product (KIAA0649), mRNA	Homo sapiens Misshapen/NIK-related kinase (MINK), mRNA	Homo sapiens Misshapen/NIK-related khase (MINK), mRNA	Homo sapiens origin recognition complex, subunit 5 (yeast homolog)-like (ORCSL) mRNA, and translated products	Homo sapiens origin recognition complex, subunit 5 (yeast homolog)-like (ORCEL) mRNA, and translated	products	Hamp sepiens chromosome 21 segment MS21C101	H.sapiens pseudogene for the low affinity IL-8 receptor	Novel human gene mapping to chamosome 1	Homo sapiens histone deacetylase 8 (HDAC8 gene) (HSAZ71724), mKNA	Homo sapiens HLA-5 gene to numen reuccoye environ b	Homo sapiens ALA-b gene for numer reaccoys a myon b
	Top Hit Database Source	EST HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN		EST_HUMAN			<u></u>	_ <b>_</b>			EST HUMAN		EST HUMAN	EST HUMAN	1	E		<u></u>	)  E		1	NT	NT	N	11	Z	<u> </u>
3		<u> </u>	<u> </u>		S	N BI	S	Ż	뉟	37 N	<u>Z</u>	8		<u> </u>	┞	ŭ	<u> </u>	N 08	8	34 N	34 N	TIM		24 N	Z	Z	Z	N 88		4
Pignio	Top Hit Acession No.	A1924663.1	BE084410.1	5.0E-66 BE898644.1	5.0E-66 BE898644.1	6679816 NT	4.0E-66 AW897798.1	X89211.1	4.0E-66 AJ223364.1	9835487 NT	4502098 NT	TN goncoak	17074	N55323.1	1	N55323.1	N55323.1		7682223 NT					4505524 NT	AL163301.2	X65859.1	AL117233.1	8923768 NT		AJ133267.2
	Most Similar (Top) Hit BLAST E Vaiue	6.0E-68 Ai	6.0E-66	5.0E-88	5.0E-66	4.0E-66	4.0E-88	4.0E-86	4.0E-66	4.0E-66	3.0E-86	9 10	3.05-90	3.05-68		3.0E-86	3.05-68	3.0E-66	3.0E-66	2.0E-88	2.0E-66		Z.UE-90	2.0E-86	2.0E-86			L		2.0E-66
	Expression Signal	1.02	1.86	0.81	0.81	2.44	1.24	2.12	2.81	4.36	34.87	60.70	34.07	102		1.02	100	2.42	7.45				1.14	1.14					39.41	39.41
	ORF SEQ ID NO:	14278	11397		14896	10833	L	12316			11455		1498	12020		12021							10061	10062			L			14496
	Econ SEQ ID NO:	720	6348	9916	9916	5803	8693	7194	7374	9623	8380		6388	6022	7	6922	6003	7595	a de la	5433	243	5	5077	5077	6782	7924	8684	9132	9511	9511
	Probe SEQ ID S NO:	4240	1349	4939	4939	782	1698	2217	2403	4638			1402	4036	2	1836	Š	3635	200	3 6	3 2	78	4	410	į	30,5	2870	4137	4521	4521

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	601508376F1 NIH MGC 71 Hamo sepiene cDNA clane IMAGE:3908831 5"	AV717817 DCB Hamo septems cDNA clane DCBADC07 6	AV717817 DCB Hamp saplens cDNA clane DCBADC07 5'	AV717817 DCB Homo septems cDNA clone DCBADC07 61	AV717817 DCB Hamo saplens cDNA clone DCBADC07 5'	BUT6002.x1 Schneider fetal brain 00004 Homo septens cDNA clone IMAGE:2782083 3' strailer to gb:M37104 ATP SYNTHASE COUPLING FACTOR 6, MITOCHONDRIAL PRECURSOR (HUMAN);	EST98812 Testis I Homo sepiens cDNA & end similar to similar to C. elegans hypothetical protein, cosmid 2X353	2156b05,r1 Sogres fetal liver spleen 1NFLS S1 Homo sepiens cDNA clone IMAGE:416049 5	THERPOR A Science faired liver spleen 1NFLS S1 Homo septens CDNA clone IMAGE:416049 6	Linea cadina incella 4 3 4 thickneshate 5/8 kinasa (ITPK1) mRNA	TOTIO Edution included by the company of the compan	Home septens incetted 1,3,4-triphosphate 5/6 kinase (11FK1), mikina	au75d02.x1 Schneider fetal brein 00004 Homo septens CDNA clone IMAGE:2782083 \$ smiler to go:M3.7104 ATP SYNTHASE COUPLING FACTOR 6, MITOCHONDRIAL PRECURSOR (HUMAN);	H.sepiens mRNA for ecestyl-CoA carboxylase	Homo saplens mRNA for transmebrane receptor protein	Homo sapiens PMP69 gene, exons 3,4,5,6 & 7	Homo sapiens retinoblastoma 1 (including osteosercoma) (RB1) mRNA			Homo sapiens DKFZp434P211 protein (DKFZP434P211), mKNA		3 (isopepudase 1-5) (USF13) mrvA			Homo sapiens T cell receptor beta rocus, I CADY/33AZ to I CADA/123A region	yn02d11,r1 Soares adult brain N2b4Hb557 Homo Baptens dDNA done invade107203 U	EST37803 Embryo, 9 week Hamo septens cLINA b end	RC4-BT0311-141189-011-h06 BT0311 Homo seprens culvA	MR3-SN0066-040500-008-f01 SN0066 Hamo septens culvA	hw16g09.x1 NCI_CGAP_Lu24 Homo sepiens cDNA done IMAGE:3183136 3' similar to WP:PZAH11.9 CE09617;	QV4-ST0234-181199-037-405 ST0234 Hamo explans cDNA	
Top Hit Database Source	EST_HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	TOT IN	EST HUMAN	EST LIMAN	NOWOL 1		<u></u>	EST HUMAN	_					T	П		_	Į,	Ę	N	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	ST HUMAN	EST HUMAN	
Top Hit Acession No.	1.0E-66 BE887173.1 ES								Ī	070230	100/243 NI	7857243 NT	4W162232.1 E				4506434 NT	4507332 NT	4507332 NT	7657020 NT	7857020 NT	7848	AF016898.1 N	AF016898.1	.1		AA333768.1		AW869159.1	BE348354.1	AW816405.1	
Most Similar (Top) Hit BLAST E Value	1.0E-88B	1.0E-88	1.0E-66	1.0E-66/A	1.0E-66/A	7.0E-87	200	<u> </u>		-1	7.0E-67	7.0E-67	7.05-67	- 1	4:3	15		8.0E-67	6.0E-67	6.0E-67	8.0E-67	6.0E-67	6.0E-67	6.0E-67	5.0E-67	4.0E-67		3.0E-67	3.0E-67	2.05-67	2.0E-67	
Expression Signal	1.17	1.49	1.49	3.88	3.88	4.85	35	108	2 3	3	1.31	1.31	4.77	153	236	1.35	1 52	1.44		3.45		0.86	1.01	1.01	202	1.2	1,65	1.01	3.04			
ORF SEQ ID NO:		12863	Ł			<u>L</u>		11410				12073	40474	L			L	L		L			14958	14959		11354			L			
SEQ ID	6840	7844	7844	7844	7844	5454	1	0000				6967	5454		SROR		1			L	L		8983	9983			L	L	L	<u> </u>	2020	Ì
Probe SEQ ID NO:	1844	2823	2823	42RO	4280	379		2021	300	1523	1982	1982	2778	2/30 EGE	787	1264	2005	3353	3353	4567	4567	4805	5012	5012	3150	1308	2741	3374	ARRE	\$	8 8	á

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Single Extensives Expressed in HBL100 Cells	Top Hit Descriptor	Homo septens manipolome (distributed in bedeeped 1)	OVASTAZA 484480 637 85 STRONG 1	Homo contone mDNA & VIA AGETT	Homo contains m DNA 4-VIA Agent	60117200251 NILL MCC 47 L	801177002F1 NIH MCC 17 LOTTO Suplets CUNA clone IMAGE:3632344 5	Homo captains and Blood Colombia applieds control of the Colombia and	Homo carione are D and make the Company of the Comp	Homo septems 282 professional and the control of th	Home earlone 386 and a secondaried page increased (POH1) mRNA	Homo sariens any less entires Search (Search Port) mRNA	Company indicate mingral by 100 (SP 100) MRNA	wm28k4 w NC COAD 11/4 U	SOLITORATION NO. 1011 TO 1011	Homo society Smad, and OV Lt	rong equals chiab- and cil-matacting zinc inger protein mRNA, partial cds	yd08a02.71 Soures infant brain 1NIB Homo septems CDNA clone IMAGE:24880 5' similer to SP:A48836	Homo septembria mRNA for KIA 44344	WH8808 XT NCT CGAP KI414 Home makes a Not Act and The Company of t	Homo saniens KIAA0553 amien cene complete color and albumin	Homo sapiens KIAA0553 protein gene complete cds; and schillin protein gene, partiel cds	Homo sapiens KIAA0553 protein gene complete cds: and sinhally and a partial cds.	Homo saplens KIAA0553 protein gene, complete cds. and alphallh protein gene, partial cds.	601109444F1 NIH MGC 16 Homo sepiens CDNA clone IMAGE 3250724 F1	ZW71g02.r1 Soares testis NHT Homo saplens cDNA clone IMAGE:781682 F	Homo saplens glutamate receptor, metabotropic 8 (GRM8) mRNA	Homo saplens glutamate receptor, metabotropic 8 (GRMB) mRNA	Rattus novegious brain specific cortactin-binding protein CRDon mBNA section	nc13d12.r1 NCI_CGAP_Pr1 Hamo sapiens cDNA clone IMAGE-1008023	Homo sapiens DGS-I mRNA, 3' end	tm89f01.x1 NCI_CGAP_Bm26 Homo saplens cDNA clone IMAGE:2185306.3	tm89f01.x1 NCI_CGAP_Bm25 Homo sapiens cDNA clone IMAGE:2165305 3'	z15h04.r1 NCI_CGAP_GCB1 Homo sepiens cDNA clone IMAGE:713239 6'
XOII PIODES	Top Hit Database Source	þ	FST HIMAN	TN	TN	EST HIMAN	EST HIMAN	)   		:  -	Ŀ	-	Ę	EST HIMAN	EST HIMAN	NT TN		FST LINAN		EST HUMAN	Į.	N	N	N	EST_HUMAN	Г	Г	N	L'N	EST_HUMAN	Г	EST_HUMAN		EST HUMAN
	Top Hit Acession No.	4505222 NT	AW816405.1	AB011149.1	\B011149.1	E2980321	E296032.1	5031978	5031978 NT	5031980 NT	5031980 NT	4507184INT	123774	873830 1	I	T	T	80514.1	B037732.1	T		Γ.			2.0E-69 BE257857.1 E		4504148 N	4504148 N		4230303.1				
	Most Similar (Top) Hit BLAST E Vedue	1.0E-68	1.0E-68	1.0E-68	1.0E-68/	1.0E-68 B	1.0E-68	9.0E-69	9.0E-69	9.0E-89	9.0E-89	9.0E-69	8.0E-69.A	4.0E-69	3.0E-89	3.0E-69		3.05-89	3.0E-89	3.0E-69/	2.0E-69	2.0E-69 /	2.0E-69 /	2.0E-69/	2.0E-69	2.0E-69 /	2.0E-69	2.0E-69	1.0E-69	8.0E-70	8.0E-70 L7	7.0E-70	7.0E-70	7.0E-70 A
	Expression Signal	0.94	11.12	1.74	1.74	26.0	1.18	7.87	7.87	2.59	2.59	9.0	1.28	0.99	5.07	2.03		8.	0.97	1.66	2.01	2.01	6.22	5.22	2.69	3.44	1.2	1.2	1.7	1.55	2	4.59	4.59	237
	ORF SEQ ID NO:	10168	10364	12289		13901	14849	10083	10084	11050	11051	14977			10478	10636	-		14771	13954	10457	10458	10457	10458	11926		14924	14925	11734	12363	14226	11860	11861	1969
	ᄪᄦ			7169				5100		6021	6021	10006	8313	5551	5460	5635	-	6521	9788	8962	5438	5438	6438	5438	6835	7791	8947	9947	0999	1,66	9242	6769	62/69	6879
	Probe SEQ (D NO:	8	82	2190	2180	3903	4901	8	20	1011	1011	5035	3302	518	385	809		1524	4804	4971	129	129	402	402	<b>18</b> 6	2770	4970	4970	1684	22	4248	1	#	1890

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Homo sapiens fumor euppressor defeted in oral cancer-related 1 (DOC-1R) mRNA	Homo sapiens adenylate cyclese 3 (ADCYS) mRNA	Homo septens emyloid beta (A4) precursor protein (protesse neidn-II, Alzheimer disease) (APP), mRNA	Human Ku (p70/p80) subunit mRNA, complete cds	Homo sepiens CMP-N-spetylneuraminic acid synthese (LOC53907), mRNA	Homo sapiens KIAA0792 gene product (KIAA0792), mRNA	Homo sapiens KIAA0792 gene product (KIAA0792), mRNA	RC0-BT0522-071299-011-e12 BT0522 Hamo septens cDNA	RC0-BT0522-071289-011-e12 BT0522 Homo septens cDNA	Homo saplens phosphatidylinosital 4-kinase 230 (p/4K230) mRNA, complete cds	y07a10.r1 Soures melanocyte 2NtHM Homo sapiens cDNA clone IMAGE-270522 6' struiter to SW:D3HI_RAT P29288 3-HYDROXYISOBUTYRATE DEHYDROGENASE PRECURSOR:	yy07a10.r1 Soares melanocyte ZNbHM Homo sapiens CDNA clone IMAGE:270522 6' similar to SW:D3HI RAT P28266 3-HYDROXYISOBUTYRATE DEHYDROGENASE PRECLIRSOR	qx51h01x1 NCI_CGAP_Pan1 Homo sapiens cDNA clone IMAGE:2004913 3'	Homo sapiens hypothetical protein FLJ20758 (FLJ20758), mRNA	Homo saplens KIAA0193 gene product (KIAA0193), mRNA	Homo sapians KIAA0183 gene product (KIAA0183), mRNA	2945h05.r1 Stratagene HeLa cell s3 837216 Homo sapiens cDNA clone IMAGE:812441 6' similar to TR:G1041283 G1041283 D2085.5;	2945h05.r1 Stratagene HeLa cell s3 837216 Homo saplens cDNA clone IMAGE:612441 6' similar to TR:G1041283 G1041283 D2085.5:	Homo sepiens chromosome 21 segment HS21C002	248g04.r1 Scares retine N2b4HR Homo septens cDNA clone IMAGE:380214 5' stratiar to SW:GAG_HTL1A Prosast GAG POI YOROTEIN	Homo saplens mRNA for KIAA0601 protein, partial cds	Novel human gene mapping to chomosome X	Homo saplens Spast gene for spastin protein	Human nonmuscle myosin heavy chain-B (MYH10) mRNA, partial cds	Homo saplens ADP/ATP carrier protein (ANT-2) gene, complets cds	Homo saplens ADP/ATP carrier protein (ANT-2) gene, complete cds	Homo esplens transglutaminese 3 (E polypeptide, protein-glutamine-gamme-glutamytramaferase) (TGM3) mRNA
Top Hit Defabase Source	¥	M	- <u>F</u>	IN	IN	IN IN	TN	EST_HUMAN	EST_HUMAN	LN	EST_HUMAN	EST HUMAN	EST HUMAN	L	IN	IN	EST_HUMAN	EST HUMAN	N.	TAT HIMAN	LZ	TN	Ę	FZ	TN	TN	– ₽
Top Hit Acesslon No.	5031668 NT	4757723 NT	4602168 NT	M30838.1	R923899 NT	7662307 NT	7662307 NT	BE071798.1	BE071796.1	AF012872.1	N42161.1	N42161.1	AI246899.1		7661983 NT	7681983 NT	AA180093.1	2.0E-70 AA180083.1	2.0E-70 AL163202.2	A A O 5 40 10 1	AB011173.1	AL133207.2	AJ246003.1	M69181.1	L78810.1	L78810.1	4507478 NT
Most Similar (Top) Hit BLAST E Value	7.0E-70	7.0E-70	6.0E-70	6.0E-70	6.0E-70	6.0E-70	5.0E-70	3.0E-70 B	3.0E-70 B	2.0E-70 A	2.0E-70 N	2.0E-70 N	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70	2.0E-70	1.0E-70
Expression Signal	6.81	3.55	4.08	4.32	1.12	1.53	1.53	3.24	3.24	1.16	11.75	11.75	1.73	233	1.5	1.5	1.22	12	1.85	7 98	1.57	1.26	1.17	4.94	1.03	1.03	3.18
ORF SEQ ID NO:		14083	10920	12169	12525	12561		11610	11611	10107	10714	10715		11045		11203	11678	11679			12506		13847	13928	14051	14052	
Exan SEQ ID NO:	6994		5879			7772	7772	6551	6551	5119	5705	5705		6015		6168	8612	6612					8840		6906	E906	8320
Probe SEQ ID NO:	2011	4103	098	2078	2437	2481	2481	1554	1554	39	080	089	695	1005	1165	1165	1615	1815	1703	2756	2415	3728	3838	3936	4069	4069	3309

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	Top Hit Desorbtor	Home serviews SP400-HMG nuclear autoantigen (SP100) mRNA, complete cds	CANA CITIZA A 44 A CO A CATANIS STD234 Homo sablens CDNA	CV4-31 UZ34-161163-001-100 Clinard) substraitly, member 10 (TNFSF10) mRNA	Hamo septents united floorest from the first of the following the first of the firs	Equits capanus grycararum year processes mental cus	Equus Garantis gives account of the control of the	Homo sapiens pasminogen (PLO) innova	Homo sapiens SP 100-mm increase secretary (SOUL), mRNA	Homo sprens promosome 21 segment HS21C008	Titulio septicili oriente de la conservant filmablests. NDHSF Homo septicins CDNA clone IMAGE:1665916 3' efinifier to	contains LOR1 b2 LOR1 repetitive element;	Homo sapiens neurona del dearry existe proven (1000) // 1000 /	Homo sapiens disabled 2 garle, tawns 2 unicupin to the complete cds	Homo sapiens priospriaudyni volus	Homo sapiens PMS2L16 mkNA, partial cas	Homo sapiens PMS2.16 mRNA, partial cds	Homo sapiens hairy/enhancer-of-split related with TKFW mouthly (TIE: L.), III C.	Homo sapiens inorganic pyrophosphatase mRNA, complete cds	sp.		ated Transcripts Hamo suplens aDNA	clone 02 15 5 similar to Homo septens chromosome 19	02_15 Human Epidemina Negalino Common 19			And Spares Infant brain 1NIB Homo saplens cDNA ctone IMAGE: 52628 6	MINOSTRIA A NCI CGAP Lu19 Homo septens cDNA clone IMAGE:2423188 3' similar to TR:088705 085709	HYPOTHETICAL 38.6 KD PROTEIN, contains Alu repetitive element,			Homo sapiens acontrase 2, mitochondrial (ACO2), nuclear gene encoding mitocondrial process, university and sapiens acontrase 2, mitochondrial (ACO2), nuclear gene encoding mitocondrial process, university and sapiens acontrase 2, mitochondrial (ACO2), nuclear gene encoding mitocondrial process, university and sapiens acontrase 2, mitochondrial (ACO2), nuclear gene encoding mitocondrial process, university and sapiens acontrase 2, mitochondrial (ACO2), nuclear gene encoding mitocondrial process, university and sapiens acontrase 2, mitochondrial (ACO2), nuclear gene encoding mitocondrial process, university and sapiens acontrase 2, mitochondrial (ACO2), nuclear gene encoding mitocondrial process, university and sapiens acontrase 2, mitochondrial (ACO2), nuclear gene encoding mitocondrial process, university and sapiens acontrase 2, mitochondrial (ACO2), nuclear gene encoding mitochondrial process, university and sapiens acontrase 2, mitochondrial process, university and sapiens acontrase 3, mitochondrial process, university acontrase 3, mitochon	
-	Top Hit Database Source		T	EST HUMAN				_				EST_HUMAN									=		EST_HUMAN	CST HIMAN	1	1	14441111	EST HOMAN	EST_HUMAN		EST HUMAN	- <b>5</b>	
		1	zΪ		뒬	뉠	뉠	틸	뉠	킰	킼	<u> </u>	털	뉟	<u>눌</u>	E	Ż	2	2 2	4	<u> </u>	4	Ü	<u> </u>		+	=	╬	·	H	╣	<del>- 2</del>	1
6	Top Hit Acession No.		5.0E-71 AF058322.1	¥8.16 <u>¥</u>	4507592 NT	4.0E-71 AF157628.1	F157628.1	4505880 NT	4.0E-71 AF056322.1	7657602 NT	L163206.2	1077927.1	7706281 NT	1.0E-71 AF205890.1	1.0E-71 AF012872.1	1 0F.71 AB017007.1	4 DE 74 AB017007.1	TRETARA NT		1F11B665.1	1.0E-71 AF246219.1	AF246219.1	BE122850.1	100000	A 504900.1	AFZIGSU4.1		H23176.1	A1857635.1		A1857635.1	4501866INT	
ŀ		$\dashv$	딁	71 A	7.1	71 A	71 A	17.	71 A	년 기	2.0E-71 AL	1.0E-71	F	71/	1	Ī,	Ī	†	Ę	1.0E-71	Ì	1.0E-71	1.0E-71	7	1.00-17	1.01-73	1.0E-71	1.0E-71	D 0F-72		9.0E-72	7 OE-73	
	Most Similar (Top) Hit BLAST E Value		5.0E-	5.0E-71 AV	4.0E-71	4.0E-	4.0E-	4.0E-71	4.0E	4.0E-71	2.0E	L														1							
	Expression Signal		14.02	0.87	76.0	215.31	215.31	1.81	7.98	6.12	16.61	1.87		4.33								5.34	0.73				9 2.05	1.19	72.0		1 0.74	78 7	
	ORF SEQ ID NO:		12244	13986	19191	10409	10410	12851	14282	14792	11250		10978						12693	13456	13534	13535	13575			13667	14319	<u> </u>		10400	10461		138/1
İ	Exon SEQ ID NO:		7127	8997	5180	5400	5400	7836	9286	9811	6210	SARO	5044	9090		2013	7012	7012	7583	8430	8524	8524	0570	3	8570	8662	9336	9450		544	74		8992
	Probe SEO ID SEO ID		2148	1004	103	348	a a a	2848	4304	4827	1210	ŝ	3 5	17007	3	1320	888	2029	2621	3422	3516	3516	6000	365	3563	3656	4345	4460		<del>6</del>	405		3996

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Single Exon Probes Expressed in HBL100 Cells

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Silving Example Topics and the second of the	Top Hit Descriptor	Homo sapiens accritisse 2, mitochondrial (ACO2), ructeer gene encoding mitocondrial protein, mRNA	Homo sapiens aconitase 2, mitochondrial (ACO2), nuclear gene encoding mitocondrial protein, mRNA	QV0-CS0010-150800-398-611 CSU010 Hand Septemble CNVA	QV0-CS0010-150800-398-611 CS0010 Italia espirale COVA	QV0-CS001U-15080U-S80-e11 CS0010 mails depicts const.	CVU-COUNTY-TOWN-COUNTY-TOWN-COUNTY-CO	From Septems alpha mindella Interval Composition (17057820.2), mRNA	TOTIC SEPTEMBLY IN THE PROPERTY OF THE CASE	HOTO SECIETS MIXAN IN NAME IN COLOR PROPERTY PRINCE OF THE	Homo explens pre-to-dell conditionally leaded (L.C.L.) his decision of the conditions of the condition	and allows I some estimate the property of the source of t	Human chondroitin sulfate proteogrycan versican V0 splice-warlent precursor peptide mRNA, complete cds	Human chondroitin suifate proteoglycan versican VO splice-variant precursor peptide mRNA, complete cds	Human gamma-aminobutyric acid transaminase mRNA, partai cds	Human gamma-amilinobulying action transcription of the control of	Hamo sapients sus no canny beneat the suspension of the suspension	Tropy vietta 2.C sinha = T-cell receptor delta and C sipha fusion gene (alternatively spliced, splice junction)	[human, precursor B-cell line REH, mRNA Partiel, 211 nt]	Homo sapiens hyponieural process (LD 11121),	Home septens protein methythensferase (JBP1) mRNA, complete cds	instance 1 Spares parethyroid tumor NbHPA Home sapiens cDNA clone IMAGE:1387395 3'	MRA CT0063-071099-002-h11 CT0063 Home saplens cDNA	WASSEGB X1 NCI CGAP BITZS Hano sapiens cDNA clone IMAGE:2501098 3' similar to TR:Q50050	Q59050 HYPOTHETICAL PROTEIN MJ1656.;	Homo sapiens hypothetical protein FLIZUSUS (FLIZUSUS), IIII NO.	Homo sapiens chromosome 21 segment HSZ10000	Homo saplens chromosome 21 segment HS210382	Homo sapiens chromosome 21 segment HS21C018	
1 110000	Top Hit Databese Source		<u> </u>	EST_HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN	IN.		¥	E	EST HUMAN	뒫	<u>₹</u>	¥	Į	5	<u> </u>	Į.	Ę	LZ!	TN TOT	TOTAL HOMAN	ESI JONESI	EST HUMAN	TN	K	IN	NT	
Single	Top Hit Acessian No.	4501888 NT	4501868 NT	333707.1	-333707.1	-333707.1	F333707.1	L11645.1	11034844	4.0E-72 AB033104.1	1976	3.0E-72 AA723823.1	3.0E-72 U16306.1	116306.1	J80226.1	J80228.1	AJ22904	8923548 NT	S77589.1	11416196	AF167572.1	AF167572.1		3 AW374968.1	3 AW071755.1		Al 16320			200
	Most Similar (Top) Hit BLAST E Value	7.0E-72	7.0E-72	6.0E-72 BF	5.0E-72 BF	5.0E-72	6.0E-72 BI	5.0E-72 L1	4.0E-72	4.0E-72	3.0E-72	3.0E-72	3.0E-72	3.05-72	3.0E-72			3.0E-72	3.0E-72					9.0E-73	8.0E-73					
	Expression	1.6	. 20.	2.81	2.81	18.08	18.08	2.73	1.32	1.07	2.48	1.27	11.37	44 97			12.79	2.5	3.01		1.08			1.26	227				\$   \$	<u> </u>
	ORF SEQ ID NO:	13978	13979			10148	10149			14988			11171	7,44		L		13242	13734		14593	14594	5 12110	0 11486	44080			13204		2
	Exan SEQ ID NO:	8992	<u> </u>			L		6128	9661	L	L		l	_	6177		١		8736		L		L	3 6430	0000		1			7 6223
ļ	Probe SEQ ID NO:	3886	9008	8	2	88	95	1122	4676	6048	19	892	1137		1137	1174	3001	3205	2722	4414	4621	4621	202	1433		1021		3227	4787	157

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Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	CARD-CN0044-280100-164-f08 CN0044 Hamo septens cDNA	Hamo sepiens heme-binding protein (HEBP), mRNA	Homo sepiens heme-binding protein (HEBP), mRNA	Homo explens BASS1 (BASS1) mRNA, partial cds	RC3-NN0066-270400-011-c04 NN0066 Homo sapiens cDNA	Homo sapiens caspase 8, apoptosis-related cysteine protease (CASP8) mRNA	Homo sepiens chromosome 21 segment HS21C083	AU121585 MAMMA1 Homo sapiens cDNA clone MAMMA1000490 5	Gallus gallus Dach2 protein (Dach2) mRNA, complete cds	Homo sapiens CD39-like 4 (CD38L4) mRNA	Homo sapiens NKG2D gene, excn 10	Homo sepiens chromosome 21 segment HS21C046	Homo sapiens S164 gene, partial cds; PS1 and hypothetical protein genes, complete cds; and S171 gene, nartial cds	xn78g07.x1 Soares_NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE:2700636 3'	601283521F1 NIH_MGC_44 Homo septens cDNA clone IMAGE:3605463 6	601283521F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3805453 5	UI-H-BIO-eah-h-03-0-UI.s1 NCI_CGAP_Sub1 Homo sapiens cDNA clone IMAGE:2706365 3	JI-H-BIG-eah-h-03-0-Ui.s1 NCI_CGAP_Sub1 Homo sapiens cDNA clone IMAGE:Z/08355 3	hr54e11.x1 NCI_CGAP_Kid11 Homo sapiens cDNA cione iMAGE:3132332 3	hr54e11.x1 NCI_CGAP_Kid11 Homo sepiers cuiva cione invace:3132332 3	Homo sapiens DEAD/H (Asp-Gill-Alia-Asp/His) box polypepide 11 (3.09 evisius Chi. 1-110 inspense) (DDX11) mRNA	Homo sapiens DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 11 (S.cerewisias CHL1-like helicase) (DDX11) mRNA	df17c09.y1 Morton Fetal Cochiea Homo sepiens dDNA clone IMAGE:2483704 67	PM0-CT0289-271089-001-h07 CT0289 Homo sapiens cDNA	Homo saplans DNA for emyloid precursor protein, complete cds	Homo sapiens mRNA for KIAA1019 protein, partial cds	Home sapiens DNA, DLEC1 to ORCTL4 gene region, section 1/2 (DLEC1, ORCTL3, ORCTL4 genes,	complete cds)	Homo sapiens DNA, DLECT to OKCILA gene region, second 1/2 (DLECT, OKCILA, OKCILA, Select, complete cds)	Homo sapiens protessome (prosome, macropain) subunit, beta type, 1 (PSMB1) mRNA	
20001110	Top Hit Database Source	EST_HUMAN				EST_HUMAN			T HUMAN	NT	17	NT	TN	Ę	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	— <b>È</b>	<u>_</u>	EST HUMAN	EST HUMAN	Į.	Z		NT.	<b>⊑</b>	Į.	
	rop Hit Acession No.	W843789.1	11435913 NT	11435913 NT	2.0E-73 AF139897.1	2.0E-73 AW898081.1 E	2682		U121585.1	F198349.1	4557428 NT			E100007 4	W263177.1		E388260.1		4W014039.1		BE048846.1	4768135 NT	4768135 NT	AWOZOBB 1	AW387756 1	Γ			AB026898.1	AB026898.1	4506192	
	Moet Similar (Top) Hit BLAST E Value	3.0E-73 A	3.0E-73	3.0E-73	2.0E-73	2.0E-73	2.0E-73	2.0E-73	1.0E-73	1.0E-73 A	8.0E-74	7.0E-74	7.0E-74	R OF 74 A	8.0E-74	6.0E-74 B	8.0E-74	8.0E-74	6.0E-74	6.0E-74	6.0E-74	6.0E-74	A 0E-74	5.0E-74	R 0E-74	4 0F-74	4 OF 74		4.0E-74	4.0E-74	4 0F-74	
	Expression Signed	88.	7.	1.1	237	3.12	3.99	1.02	2.61	1.04	1.76	2.57	1.22	-	0.92	63.86	53.86	0.97	76.0	1.37	1.37	1.65	4	1.50	A 54	7 A	2	3	1.42	C7 1	R 12	
	ORF SEQ ID NO:	11367	11909	11910	10900		13143		11819	12503	10775	11989	13288			12347	12348		12832		13643	14821	<u> </u>	4822		0,200			12002	_	1	
	Exam SEQ ID NO:	6319	6815	6815	2860	6893	8124	9303	6740	7384	5754	6895	8266	250	8584	7229	7229	7813	7813	8637	8637	0848	1 _			_	2555	1	6908	1	1	i
	Probe SEQ ID NO:	1321	1825	1825	2	1906	3108	4311	1745	2413	731	1909	3253	7,40	1587	2252	2252	2793	2793	3631	3631	4866		888		7887	7/2	242	1922	1	7 67	2

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Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo sapiens proteasome (prosome, macropain) subunit, beta type, 1 (PSMB1) mRNA	Homo septems mRNA for KIAA1168 protein, pertial cds	Homo sapiens PLP gene	Homo seplens chromosome 21 segment HS21C010	Homo sapiens chromosome 21 segment HS210047	Hamo sapiens KIAA0569 gene product (KIAA0569), mRNA	Homo sapiens mRNA for transmebrene receptor protein	Homo sapiens hydroxyscyl-Coenzyme A dehydrogenase/3-katoacyl-Coenzyme A thiolass/encyl-Coenzyme A hydratase (trifunctional protein), beta subunit (HADHB) mRNA	Homo sapiens hydroxyscyl-Coenzyme A dehydrogensse/3-ketoscyl-Coenzyme A thiolesse/enoyl-Coenzyme A hydratese (trifunctional protein), beta subunit (HADHB) mRNA	Homo saplens mRNA for KIAA1442 protein, partial cds	Homo saplens glyceraldehyde-3-phosphata dehydrogenase (GAPD), mRNA	Homo sapiens glyceraldehyde-3-phosphate dehydrogenase (GAPD), mRNA	Human endogenous retrovirus HERV-K-147D	wx51e07.x1 NCI_CGAP_Lu28 Homo septens cDNA clone IMAGE:2547204 3' similar to SW:GG95_HUMAN_CORRATO COI CIN DS revertains element MFR22 manifilitie element :		Homo septems epidemma growm ractor receptor (awan crymrobastor reuncama was (v-cru-u) croopere homolog) (EGFR) mRNA	Homo sapiens epidermal growth factor receptor (avian erythroblastic leukernia viral (v⊷erb-b) oncogene homolog) (EGFR) mRNA	PT2.1_15_G11.r tumor2 Homo saplens cDNA 3'	Novel human gene mapping to chomosome 22	Novel human gene mapping to chomosome 22	Human platelet glycoprotein IIb mRNA, 3' end	Homo sapiens Mischapen/NIK-related kinase (MINK), mRNA	QV4-ST0234-181199-037-f05 ST0234 Homo sapiens cDNA	Homo sapiens hypothetical protein FLJ11026 (FLJ11026), mRNA	Homo saplens beta 2 gene	Homo saplens zinc finger protein 259 (ZNF259) mRNA	Homo saplans chromosome 21 segment H3210346	Homo sapiens DNA for Human PZXM, complete dos	Home sapers mannessasse, aprile, was 27, mannes 1 (mm 27.1), mm 4.7.	Light o sphore grant and the control of the control
	Top Hit Database Source														TOT LINAMIN	- TOWER			EST HUMAN					EST HUMAN							
L		Σ	MI	호	Z	눌	호	둗		Ē	上	볼	볻	ž	- 6	2	៑	<u> </u>		ż	ż	Ż	Z	Ĕ	N 6	Ż	Z	z	회		딁
	Top Hit Acession No.	4508192 NT	4B032994.1	4.0E-74 AJ006976.1	4.0E-74 AL163210.2	4.0E-74 AL163247.2	7862183 NT	217227.1	4504326 NT	4504328 NT	B03786	7669491 NT	7669491 NT	AF020092.1	7 000000	Z.UE-/4 AI8505Z6.1	4885198 NT		A1557280			J02963.1	TR57334 NT	AW8164		X02344		AL1632	AB00205		1 NI 01 1 1 0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
	Most Similer (Top) Hit BLAST E Velue	4.0E-74	4.0E-74	4.0E-74	4.0E-74	4.0E-74	4.0E-74	4.0E-74 Z	4.0E-74	4.0E-74	4.0E-74 A	2.0E-74	2.0E-74	2.0E-74 A	100	Z.UE-/4	2.0E-74	2 0F-74	2.0E-74 A	2.0E-74	2.0E-74 /	2.0E-74	1.0E-74	1.0E-74		1.0E-74	1.0E-74	1.0E-74			1.0E-74
	Expression Signal	6.12	1.18	6.03	0.81	122	1.98	1.19	1.02	1.02	0.81	263.61	263.61	12		2.76	3.33	2 83	3.89	2.77	2.77	3.98	292	4.23	1.18	29.75	2.36	2.04	Ş	3.19	0.67
	ORF SEQ ID NO:	12107	12166						_					L		11270	11618							10394			L	11030			13831
	SEO ID NO:	7002	L		_		1			1		L			<u> </u>	6224	6557	<u> </u>								L	L				8824
	Probe SEO ID NO:	2019	2074	3017	3449	3950	4425	4481	4902	4902	5011	g	PA3	1156		1225	1560	4590	2528	4848	4846	4850	2	335	496	503	596	88	2165	3066	3822

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Table 4
Single Exon Probes Expressed in HBL100 Cells

																	_		<u>۲</u>	-	1	<u> </u>	¥	=	<u>.                                    </u>	~		-	<del></del>	<u>الله</u>	41.		
Top Hit Descriptor	Homo sapiens gutamate receptor, tonotropic, kalnate 1 (GRIK1) mRNA	Homo capiens chromosome 21 segment HS21 0068	hz73h08 x1 NCL_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3213663 3' similar to WP:B0611.12	Union services DNA extension-5 methydransferses 38 (DNMT38) mRNA, complete ods	FIGURE SEPTEMBER OF THE PROPERTY OF THE PROPER	WK38808.X1 NG_CGAP_PTZZ HOMD SEDING CHAN CHAN IN A CELZA 17004 S SITTER TO CGAP_PTZZ HOMD SEDING CHANNAN); RETROVIRUS-RELATED POLYPROTEIN (HUMAN);	QV1-BT0632-210200-079-602 BT0632 Homo saplens cDNA	yx30h08.r1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:249056 b	CM0-NN0057-150400-335-a11 NN0057 Hamo sapiens CUNA	601303886F1 NIH_MGC_21 Home sapiens cDNA clone IMAGE:3636344 0	Homo sepiens hypothetical protein FLJ10747 (FLJ10747), mRNA	Homo sapiens HTRA serine protease (PRSS11) gene, complete ods	Homo sapiens HTRA serine protesse (PRSS11) gene, complete cds	Homo sepiens mRNA for KIAA0581 protein, partial cds	Homo saplens platelet-derived growth factor receptor-like (PDGFRL) mKNA	Homo saplens synaptojanin 1 (SYNJ1), mRNA	Homo sapiens synaptosomal-associated protein, 28kO (SNAP29) mrtNA	Home sapiens chromosome 21 segment HS21C001	Homo septens mRNA for KIAA0581 protein, pertial cds	Human calcium-dependent phospholipid-binding protein (PLAZ) mrt/V4, campiers cas	Human calcium-dependent phospholipid-binding protein (PLAZ) minny, compress cus	Homo sapiens DNA for amyloid precursor procein, complete cas	Homo septiens KIAA0971 protein (KIAA0971), mKNA	Homo capiens chromosome 21 segment HS21C009	xg60d02.x1 NCI_CGAP_Ut4 Homo saplens cDNA done IMAGE:2332/07 3 similar to contains 1 1 17.11	PIK/ repetuve element	H. saplens ERCC2 gens, coms 1 & 2 (pertial)	601167633F1 NIH_MGC_21 Home saplens cDNA clone IMAGE:3504212 b	601437130F1 NIH_MGC_72 Home caplens cDNA clone IMAGE:3822303 b	wb30b10.x1 NCI_CGAP_GC6 Homo sapiens cDNA clone IMAGE:2307163 3' struiter to 11k:U70239 U76239	TRAP1;	WESUDION NO. CAMP GOO FAILD SQUEE CONTROLL CONTR	
Acession Top Hit Detablese Source	π	IN		ESI HOMAN		EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	7	NT	NT	NT	. 17	NT	N.	NT	NT	NT	NT	NT	NT	NT		EST HUMAN	NT	EST_HUMAN	<b>EST HUMAN</b>		EST HUMAN	EST HUMAN	_
Cop Hit Acession No.	4504116 NT	1.0E-74 AL163268.2			8.0E-75 AF176228.1	6.0E-76 AI817415.1		136757.1	4.0E-75 AW897230.1	E409464.1	8922637 NT	4F157623.1	4F167823.1	4B011153.1	5463871 NT	4507334 NT	9153	AL163201.2	AB011153.1	M72393.1	M72393.1	D87675.1	7662421 NT	AL163209.2		AW168135.1	X52221.1				AI652648.1	A1652648 1	7
Most Similar (Top) Hit BLAST E Value	1.0E-74	1.0E-74/		1.0E-74	8.0E-75/	6.0E-75	4.0E-75	4.0E-75	4.0E-75	4.0E-75B	4.0E-75	3.0E-75	3.0E-75	3.0E-75	3.0E-75	3.0E-75	3.0E-75	3.0E-75	3.0E-75	3.0E-75	3.0E-75		3.0E-75	3.0E-75		1.0E-75	1.0E-75				9.0E-76	90.20	
Expression Signal	0.67	4.87		1.12	3.52	96.0	3.67	0.99	1.27	5.17	0.97	2.28	2.29	1.84	2.26	0.82			1.01	0.75	0.75	1.58	0.82			30.85	3.35				3.62	8	
ORF SEQ ID NO:	13832			14124		12352			11801	12818						12147							L			12334			L		10116		101
SEQ ID NO:	8824	8863		946	7536	7234	5184	5483	6723	7877	8429	6002	6002	6791	6876	7035	7330	7974	8132	8285	8285	L	L			7216	L				5128		250
Probe SEO ID NO:	3822	3881		4145	2573	2257	112	458	1728	2776	3421	987	888	180	1887	2053	2356	2955	3116	3273	3273	4044	4314	4996		2239	2876	4552	1 20	285	46		\$

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo sapiens H factor 1 (complement) (HF1) mRNA	Homo saplens H factor 1 (complement) (HF1) mRNA	Homo saplens mediator (Sur2), mRNA		Homo sapiens ditydrollpoarride dettydrogensee (E3 component of pyruvate dettydrogensee complex, 2-coo- chizerate complex, branched chain keto acid dehydrogenase complex) (DLD) mRNA	Homo sapiens cAMIP-specific phosphodiesterase 8A (PDE8A) mRNA, partial cds	Homo saviens hymphocyte antigen 75 (LY75) mRNA, and translated products	1 - Consideration and unless (7 8-dilhydroblem:NADP+ and oreductase) (SPR) mRNA	Home Sapirativa in Fourtees (7.5 Animaterin NADP+ exdereductase) (SPR) mRNA	Homo sapiens seguiaposin reducioses (1,00m/parenteres)	COUNTY AND A COMPANY OF THE COUNTY OF THE CO	Human mkNA for nivides of the second	Human mRNA for Hind-1, complete cas	Human mRNA for HMG-1, complete cas	QV3-BN0047-270700-283-g06 BN0047 From septems CDNA close IMAGE:3083882 3'	UI-H-BW 1-enz-b-04-U-U.ST NCI COMP CIND INCIDENCE CON COMP INA GE-3083882 3	UI-H-BW 1-anz-b-04-C-UI-S1 NCI_COOK_ Sub/ Hallo apparis Extra Subs III NCI_COOK_ Subs III NCI_COOK_ Subs III NCI_COOK_ Subs III NCI_COOK_ Subs II NCI_COOK_	Homo sepiens eukaryodo uzusaduni edungadun isatur. 1 beta 2 (EEF1B2) mRNA	Home seprens envelyour delicency was graded from Seprens CONA	RCS-S 1030C-13010C-0.33-Au3 G 10300 Form septemb Committee Committ	Tropolicourted to the profession TPRDII complete ads	Turner miles in possible protein TPRDII complete cds	Himan mRNA for notestitle protein TPRDII, complete ods	Indiana marks for possible processing (Chroad hinding protein 1 (IGBP1) mRNA		nome suprema gravescar (COC) in the control of the	HOMO SADIERS CAWL TESTOCIONE CONTROL (CAVOA) mRNA	Homo sepiens GMZ ganglioside acuvata protein (CMZA) IIII WAS	Homo sapiens GMZ gangiloside acuvatur protein (Cwizz) III v. C	OLFACTORY RECEPTOR-LIKE PROTEIN FO	zw64e02.s1 Sceres_testis_NHT Homo septens CUNA cighe IMAGE.1 00500.5 Sulfilled CONTINUED TO THE PROPERTY PRECURSOR.;	2x64e02.s1 Soares testis NHT Homo sepiens cDNA clone IMAGE:780988 3° similar to SW:ITB6_HUMAN	P18084 INTEGRIN BETA-5 SUBUNIT PRECURSOR.;	
	Top Hit Database Source						12		-	=	Ę	EST_HUMAN	Ā	Ę	NT	EST HUMAN	EST HUMAN	EST HUMAN	Ę!	±N.	EST HUMAN	EST HUMAN	Ž!	Z	Į.	Į.	Z	LN.	N	NT	SWISSPROT	EST HIMAN		EST_HUMAN	
,	Top Hit Acession No.	4504374 NT	4504374 NT	TV ACTANT		COMPLOS	ENERGO 4	200	720c0c4	4507184 NT	7184	_		D63874.1	D63874.1			BF516282.1	4503476 NT	8476 8476		BF375689.1	D84295.1	D84295.1	084295.1					4504028 NT	P23268	44460004		AA445982.1	
	Most Similar (Top) Hit BLAST E Vatue	8.0E-78	8 0E-76	20.00	0.0E-70	9	1,05-70	1.05-70	7.0E-76	7.0E-76	_		6.0E-76	6.0E-76	5.0E-76	4.0E-76	3.0E-76	3.0E-76	3.0E-76	3.0E-76	3.0E-76	3.0E-76		2.0E-76	2.0E-76				2.0E-76	2.0E-76	L		ZUE-/0	2.0E-76	
	Expression Signal	0.84	28.0		-		8 !	74.0	7.12	4.97	4.97	18.64	15.24	15.24	15.24	0.8	1.54	1.54	21.41	21.41		6.25			1.94	1.42	1.68	1.16	2.34				1.89	1.89	ļ
	ORF SEQ ID NO:	10974	4007F	0/801	128/8					14223	14224		11983		L		10653		11623	11624		13371	10351		10399	15	3 10615	11056			l		13260	13261	
	Ean SEQ ID NO:	5041	200		<b>1859</b>		6788	8239	8241	9240	9240	6213	0689							6561		8323	5337	5392	5392	5495	5616	L	L		L	1_	3 8238	3 8238	ļ
	Probe SEQ ID NO:	700	120	474	2839		787	3220	3226	4248	4246	1214	1903	1803	1903	3134	623	623	1564	1564	3344	3344	279	340	340	458	585	1014	1504	1504	200	2	3223	3223	

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	ac83b02.y6 Stratagere lung (#837210) Homo eaplens cDNA clone IMAGE:889163 & similar to TR:014591 014591 SIMILARITY TO P22059 ;	zu/0g11.r1 Soeres_testis_NHT Homo sepiens cDNA clone IMAGE:743398 6' similer to WP:R05D3.2 CE00281;	Homo saplens chromosome 21 segment HS21C083	QV3-0T0028-220300-132-b11 OT0028 Homo sepiens cDNA	Human mRNA for HMG-1, complete cds	Human mRNA for HMG-1, complete cds	yp11h02.r1 Soeres bresst 3NbHBst Hom <b>o eaplens cDNA clone INA</b> GE:187166 6' <b>shnikar to</b> SP:ANKB_HUMAN Q01484 ANKYRIN, BRAIN VARIANT 1;	601866928F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:4109509 61	zu91g01.s1 Soeres_testis_NHT Homo sepiens cDNA clone IMAGE:745392.3'	Homo sapiens polymerase (RNA) ii (DNA directed) polypeptide E (25kD) (POLRZE) mRNA	Homo saplens polymerase (RNA) II (DNA directed) polypeptide E (25kD) (POLR2E) mRNA	Homo sapiens interferon (alpha, beta and omega) receptor 2 (IFNAR2) mRNA	qe77h12.xt Soares_fetai_lung_NbHL19W Homo sapiens cDNA clone IMAGE:1745063 3'	Homo sapiens midline 1 (Opitz/BBB syndrome) (MiD1) mRNA	Homo sapiens midline 1 (Opitz/BBB syndrome) (MID1) mRNA	7 Homo sapiens glucokinase (GCK) gene, excn 2	Homo sapiens disintegrin and metalloprotease domain 10 (ADAM10) mRNA	Home sapiens tousled-like kinase 1 (TLK1) mRNA, complete cds	Homo saplens cullin 1 (CUL1) mRNA	Homo sapians ubiquitin specific protesse 18 (USP18), mRNA	Homo sepiens EGF-like repeats and discoldin Hike domains 3 (EDIL3), mKNA	Homo sapiens EGF-like repeats and discodin Filke domains 3 (EDIL3), mixing	DKFZp434G1728_r1 434 (synonym: htes3) Hamo sapiens cDNA clara DNFZp434G1726 5	AL449758 Homo sepiens fetal brain (Stavrides GS) Homo sepiens culva	Homo sapiens SET domain and mariner transposase fusion gene (SETMAR) mRNA	Homo sapiens SET domain and mariner transposase fusion gene (SETMAR) mRNA	AV764617 MDS Homo sepiens cDNA clone MDSBTF10 5	RC3-BN0053-170200-011-h01 BN0053 Home saplens cDNA	Homo sapiens CGI-79 protein (LOC51634), mRNA	Homo saplens mRNA for KIAA1415 protein, partial cds	Homo sepiens mRNA for KIAA1415 protein, pertial cds
Top Hit Database Source	EST_HUMAN	EST_HUMAN	M	EST_HUMAN	NT	NT.	EST_HUMAN	EST_HUMAN	EST_HUMAN	TN	ΤN	TN	EST_HUMAN	TN	TN	NT	NT	N	NT.	Į,	LN T	1Z	EST HUMAN	EST HUMAN	IN	IN	EST_HUMAN	EST_HUMAN	NT	IN	Ŋ
Top Hit Acessian No.	821140.1	4400700.1		5		1.0E-76 D63874.1		8.0E-77 BF205181.1	7.0E-77 AA625755.1	4505944 NT	4505944 NT	4504600 NT	A1204088.1	4557752 NT	4557752{NT	AF041015.1	4557250 NT	AF162666.1	4503160 NT	8394518 NT	5031660 NT	5031660 NT	AL043953.1	AL449758.1	5730038 NT	5730038 NT	AV764617.1	AW997712.1	7708315 NT	AB037836.1	AB037836.1
Most Similar (Top) Hit BLAST E Value	2.0E-78 A	20E-78	2.0E-76	2.0E-76	1.0E-78	1.0E-76	8.0E-77	8.0E-77	7.0E-77	7.0E-77	7.0E-77	6.0E-77	6.0E-77	6.0E-77	6.0E-77	5.0E-77	6.0E-77	5.0E-77	5.0E-77	5.0E-77	5.0E-77	6.0E-77	5.0E-77	4.0E-77	3.0E-77	3.0E-77	2.0E-77	2.0E-77	2.0E-77	2.0E-77	2.0E-77
Expression Signal	2.	7.33	-	6.31	6.78	5.78	3.03	1.16	1.62	9.62	9.62	6.1	2.09	96.0	86.0	1.5	2.76	1.11	1.24	1.75	0.99	66.0	2.68	1.93		1.57		71.7	5.42		222
ORF SEQ ID NO:	13430				14155	14158	10281	14384	11988	12439		10329		14743		11255	11391	12689	12762		14539	14540	14749	13634				l.	12138	١	Ш
Exan SEQ ID NO:	8404	<u> </u>	L	L	9168	9168	6250	8382	L	7319	1		L	9756			6340	_	7848	_	L	9553	9764	L	L	L		6405	Ŀ	1_	Ш
Probe SEO ID NO:	3396	3682	4	4780	4173	4173	187	4391	1889	2345	2345	280	1511	4772	4772	1216	1343	2613	2690	3443	4565	4565	4780	3620	1928	1928	1334	1407	2045	2518	2518

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	ho43b05.x1 Scenes_NFL_T_GBC_S1 Hamo sepiens cDNA done IMAGE:3040113 S' similar to SW:GAG2_HUMAN P10264 RETROVIRUS-RELATED GAG POLYPROTEIN;	w22g02.x1 NCJ_CGAP_Bm52 Homo saplens cDNA clone IMAGE:2280466 3' similar to TR:066245 065245 F21E10.7 PROTEIN. ;	w22g02.x1 NCI_CGAP_Bm52 Homo sapiens cDNA done IMAGE:2260466 3' similar to TR:065245 085245 F21E10.7 PROTEIN. ;	Homo sapiens glutarnic-oxaloscetio transaminase 2, mitochondrial (aspartate eminotransferase 2) (GOT2), nuclear gene encoding mitochondrial protein, mRNA	ns68g12.s1 NCI_CGAP_P72 Homo sepiens cDNA clone IMAGE:1188839 similar to SW-RL29_HUMAN P47914 60S RIBOSOMAL PROTEIN L29. [1] ;contains element MSR1 repetitive element;	Homo sapiens mRNA for KIAA1276 protein, partial cds	Homo sapiens mRNA for KIAA1276 protein, pertial cds	Homo saplens amyloid beta (A4) precursor protein (protease nexin-il, Alzheimer disease) (APP), mRNA	Homo sapiens amyloid beta (A4) precursor protein (protease nextr-II, Alzheimer disease) (APP), mRNA	Homo saptens amyloid beta (A4) precursor protein (protesse nexin-II, Abzheimer disease) (APP), mRNA	Homo saptens amyloid beta (A4) precursor protein (protesse needn-II, Alzheimer disease) (APP), mRNA	w63605.x1 Scares, thymus_NHFTh Hamo sapients aliving additional invariances and a	Homo sapiens mRNA for KIAA1101 protein, complete cds	Homo saplens 2,4-diendy Cox regulates 1, millionarial (DEVN1), linvay	Homo sapiens CGI-60 protein (LUC-51629), mrt/V4	Home septients sold to config between niving and obtained in construction of the configuration  Homo sapiens breast cancer 1, early disert forward, a unionity state in the restriction of the contract of the	GAUGGOAT NCI_CASAP_nab namb sapatis curv cada impac	Homo sapients collegen, type Al, while I (OCLIAN), him and	Home sapiens Kirkaudus gere product (Kirkaudus), III Nama	Homo sapens NiAAuuus gene produck (NinAvus), IIII Ah	Home sapiens CAMIY Tesponsive distrant binding promit I (CALCY) in a sa	AUTIBASS HEMBAT I DOMO BENERIA COLON CINIO HEMBATON COLON CO	AUTIBIOS REMBAI notic advanta contratoria.	
Top Hit Detabase Source	EST_HUMAN	EST_HUMAN	EST_HUMAN	— <b>L</b>	EST_HUMAN	1	NT .	<b>_</b>	<b>.</b>		— <b>Ŀ</b>	EST_HUMAN	NT	F		LN	E	EST HUMAN			٦	닐	EST HUMAN	EST_HUMAN
Top Hit Acessian No.	044316.1	13519.1		304068		1.0E-77 AB033102.1 NT		4502168 NT	4502166 NT	4502168 NT	4502166 NT	·	1.0E-77 AB029024.1	4503300 NT	7706299	220	52322	1.0E-77 AI273014.1	11418424 NT	7861849 NT	7661849 NT	888	6.0E-78 AU118789.1	
Most Similar (Top) Hit BLAST E Vatue	2.0E-77 BE	2.0E-77	2.0E-77	2.0E-77	2.0E-77	1.0E-77	1.0E-77	1.0E-77	1.0E-77	1.0E-77	1.0E-77			1.0E-77	1.0E-77	1.0E-77 AJ					1.0E-77			
Expression Signal	39.	. 0.82	0.82	84	6.49	0.89	0.89	3.11	3.11	6.08	6.08	1.58		2.25				0.74	1.24	1.42	1.42	0.68		2.26
ORF SEQ ID NO:	13913				14596	10112		10340		10826		L	L	13002		14366		14527	14698	14845	14846	14341	1 10172	1 10173
Exan SEQ ID NO:	8922	9280	9280		8098		5124	6329	<u> </u>	7733	l	L	]_	7989	9221	9384	9206	9542	9713		1786	9361	5161	5161
Probe SEQ ID NO:	3922	4288	4288	4465	4623	4	4	270	270	88	88	1875	2378	2971	4227	4393	4516	4554	4728	4898	4898	4989	2	æ

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	Top Hit Descriptor	8020/8028F1 NC CGAP Brill Home series Child	Hamo seplens tracificate protein El (41348/El (41348) DAIA	ba54h03.y3 NIH_MGC_10 Homo sapiens cDNA clone IMAGE:2900405 5' similar to WP:Y48B6A.6 CE22131	Human collecteness true IV (C) G4) gene som 8	DKFZD434N03Z3 r1 434 (supervm: hises) Home carlene cDNA close DKFZ 43 A loss of H	Novel human dana manahara b ahamasama 22	Homo saciens pre-mRNA splicing factor (SERS3) mBNA Americka ad-	Homo saplens syncytin (LOC30816). mRNA	Homo sapiens phosphatidylinesital 4-kinasa, catalytic alpha polyacetida / Divaca / Divaca /	Homo sapiens phosphatidylinosital 4-kinasa, catalydia, alpha polypopula (Finatoral monta)	Homo saplens eRF1 cene. complete cde	Homo sepiens eRF1 gene, complete orts	Homo sapiens apoptasis inhibitor 3 (API3) mRNA	Homo sapiens nuclear antiden Sp100 (SP100) mRNA	AU140604 PLACE3 Homo saoiens cDNA clone PLACE3nnn273 K	Homo saplens troe IV callagen alpha 5 chain (COI 445) sere jerren 20	EST182583 Jurkat T-cells VI Homo saniens cDNA 5' and	Homo sepiens nucleoporin 155kD (NUP155) mRNA	Homo saplens peptide YY (PYY), mRNA	RC2-BN0074-090300-014-c12 BN0074 Homo septens cDNA	Homo sapiens chromosome 21 segment HS21C010	Human mRNA for KIAA0045 gene, complete cds	Human mRNA for KIAA0045 gene, complete cds	601472766T1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:3875657 3'	Homo saplens hypothetical protein FLJ10283 (FLJ10283), mRNA	Homo sapiens intersectin short isoform (ITSN) mRNA, complete cds	Homo sapiens cell-line tsA201a chloride ion current inducer protein ((Cin) gene, complete cds	Human zinc finger protein ZNF131 mRNA, partial cds	y48f03.s1 Soares fetal liver spleen 1NFLS Homo sepiens cDNA clone IMAGE:2085413'	801159415F2 NIH_MGC_53 Hamo sapiens aDNA done IMAGE:3511107 5	Hamo saplens BCL24like 2 (BCL21.2) mRNA	th18h07.x1 NCI_CGAP_Pr28 Homo saplens cDNA clone IMAGE:2118885 3'	Homo sapiens Dickkopf gene 4 (DKK-4), mRNA
	Top Hit Datebase Source	EST HUMAN	Þ	EST HUMAN	Z	EST HUMAN	NT	N	두	15	=	Į	Z FZ	  -	E	EST HUMAN	N	EST HUMAN	<u></u>	E	EST_HUMAN	LN LN	TN	IN	EST_HUMAN	11	NT	TN	NT	EST_HUMAN	EST_HUMAN	T	47.1 EST_HUMAN	<u> </u>
	Top Hit Acession No.	BF344101.1	2488	AW673424.1	Γ	Γ	4.0E-78 AL355841.1		7858876 NT	4505806 NT	4505806 NT	F095901.1	F095901.1	2142	4507164 NT	U140804.1	04489.1	_	88	11525891 NT		8.0E-79 AL163210.2			7.0E-79 BE619648.1 E	8922325 NT		(1			2.0E-79 BE379926.1  E		5237	7657024 N
	Most Similar (Top) Hit BLAST E Vatue	6.0E-78	5.0E-78	5.0E-78	5.0E-78	4.0E-78	4.0E-78	4.0E-78	4.0E-78	4.0E-78	4.0E-78	3.0E-78 A	-3.0E-78 A	3.0E-78	3.0E-78	3.0E-78 A	2.0E-78	2.0E-78	1.0E-78	9.0E-79	9.0E-79	8.0E-79	8.0E-79	8.0E-79 D28476.1	7.0E-79	4.0E-79	3.0E-79	3.0E-79	3.0E-79 U09410.1	2.0E-79 H63129.1	2.0E-79 E	2.0E-79	2.0E-79 AI	2.0E-79
	Expression Signal	0.88	1.01	4.77	4.18	1.68	1.28	20.59	1.87	1.75	1.75	3.27	3.27	1.06	1.34	1.23	2.47	1.51	1.22	3.65	5.64	0.91	1.67	1.67	19.1	0.99	1.61	5.56	2,62	1.02	1.22	1.47	1.18	0.92
	ORF SEQ ID NO:	13274	10291	12574	13339	11156	11539	12350	14168	14602	14603	10235	10236	12335	13180				14953	14533	14684	13668	14337	14338	13218		10377	11012	13051		10661	10988	1	11824
	Exan SEQ ID NO:	8253		7459	8312	6126	j		١	9613	9613	5226	5226	7217	8158	8683	8065	8907	8978	9548	6696	8663	8357	9357	8195	8119	5366	5978	8042	5343	5658	583	6028	6745
	Probe SEQ ID NO:	3240	217	2491	3301	1120	1487	224	4185	4628	4628	<u>\$</u>	160	2240	3142	3678	3048	3907	2002	4560	4714	3857	4366	4366	3179	3103	ञ्च	8	3025	285	828	915	1018	1751

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor		Homo sepiens Dickkopf gene 4 (DKK-4), mRNA	Homo eablens bhosphodiestwasse 64. cGMP-enecific md alcha (PINEAA)	Homo seplens phosphodiesterase &A. c.GMP-specific and ships (PDEst) mDNA	Homo sablens mRNA for Fas-associated factor EAE1 (Feff Ages)	Homo sapiens hepatocellular carcinoma essociated entices 88 (HCA 88) mPNA	Homo saplens mRNA for Fas-associated factor. FAF1 (Faf1 care)	ai23e05.81 Sogres testis NHT Homo sepiens cDNA clove 1349848 9	al23e05.s1 Sogres testis NHT Hamo saniens cDNA clone 1343648 3	Homo serviens V dymmesoma anomatwaesia candidata.	W4902.1 Scares placents Nb2HP Home series CNNA Alone MARCH Appears of	#58402.X1 NCL. CGAP_BM23 Homo septems cDNA clone INAGE:2103459 3' similer to SW:NUEM_HUMAN_O16785 NADH-I RICH INONE OXIONEEN ICTASE 30 KD SI ID INTO POPEM INCOME.	Homo septens NRD convertese mRNA complete ods	Homo saplens minichromosome maintenence deficient (S. comulator) 3 (MONA) DNA	Homo sabiens minichromosome maintenance deficient (S. oceanistac) 3 (MC), 13 MDNA	Homo saplens mRNA for KIAA1155 protein, partial cds	Homo sapiens mRNA for KIAA1155 protein, partial cds	Homo sapiens proteasome (prosome, macropain) 26S subunit, non-ATPassa, 3 (PSMD3) mRNA	Homo sapians serine-threonine protein kinase (MNBH) mRNA, complete cds	Homo saplens serine-threonine protein kinase (MNBH) mRNA, complete cds	H. saplens next gene (exon 12)	Homo sapiens chromosome 21 segment HS21C083	Human (3)mbt protein homolog mRNA, complete cds	Homo saplens mRNA for KIAA1434 protein, partial cds	Homo sapiens H3 histone family, member J (H3FJ) mRNA	Homo sapiens chromosome 21 segment HS21C068	Homo saplens chromosome 21 segment HS21C010	PM0-GN0018-040900-002-E03 GN0018 Homo sapiens cDNA	QV4-BN0263-040600-241-g10 BN0263 Homo sepiens cDNA	yg65a08.r1 Soares Infant brain 1NIB Homo sapiens cDNA clone IMAGE:38060 67	RET4B7 subtracted retina cDNA library Homo saptens cDNA done RET4B7	DKFZp434D1323_r1 434 (synonym: htes3) Homo saplens cDNA clone DKFZp434D1323 6
Top Hit Database Source		Į	Ę	Þ	Ė	¥	뉟	EST HUMAN	EST HUMAN	   <del> </del>	EST HUMAN	EST HUMAN	¥	Þ	Ę	LN	Ę	5	Į,	TN	IN	NT	TN	NT L	5	Z Z	41	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	EST HUMAN
Top Hit Acessian No.		7857024 NT	4585883 NT	4585863 NT	J271408.1	2.0E-79 AF244138.1	2.0E-79 AJ271408.1		9.0E-80 AA725848.1				34898.1	6631094	6631094 NT		6.0E-80 AB032981.1	6228		.1		.2		6.0E-80 AB037855.1	4504292 NT				3.0E-80 BE817465.1		2.0E-80 AI444821.1	
Most Similar (Top) Hit BLAST E	Value	2.0E-79	2.0E-79	2.0E-79	2.0E-79 A	2.0E-79	2.0E-79	9.0E-80	9.0E-80	8.0E-80	7.0E-80	6.0E-80 AI	8.0E-80 U	6.0E-80	6.0E-80	6.0E-80 A	6.0E-80	5.0E-80	5.0E-80	5.0E-80	5.0E-80	5.0E-80	6.0E-80	6.0E-80	5.0E-80	5.0E-80 AI	3.0E-80	3.0E-80	3.0E-80	2.0E-80	2.0E-80	2.0E-80
Expression Signal		0.92	3.3	3.3	2.05	2.65	1.34	18.41	18.41	1.08	1.67	2.22	2.05	4.78	4.78	1.33	1.33	5.48	1.83	1.83	1.23	1.63	1.18	8.73	6.29	1.24	15.21	1.41	7.56	3.95	1.48	3.58
ORF SEQ ID NO:			12180		12223	12346	14027	13102	13103		14784	10947	11661	12327	12328	14142	14143			10883		1	12394	12454	12792	14763		14548		11839	11907	12089
Exen SEQ ID NO:									808	8528	9781	6907	0099	7211	7211	9157			5845		6172	6426	7275	7337	7679	9780	6279	9559	9738	6754	6813	6985
Probe SEQ ID NO:		1761	2087	2087	2130	2249	4041	3073	3073	3521	4797	688	1604	2234	2234	4162	4162	<b>88</b>	825	825	1169	1429	23 23 23 23 23 23 23 24 24 24 24 24 24 24 24 24 24 24 24 24	2363	2722	4796	216	4571	4753	1762	1823	2002

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Single Exon Probes Expressed in HBL100 Cells		Top Hit Descriptor		Homo saplens chromosome 24	Homo saplens chromosome 21 unknown mRNA	INDITIZACE NCI_CGAP_Cos Homo sapleris cDNA clone IMAGE-1078466 51	2839007.1 Scenes feed in	Aku repetitive element.	2e21d10.r1 Source_fela_heart_NbHH19W Homo saplens_cONA_class_ns_con_	PEDERITION AND THE PARTY OF YOUNG TYROSINE RICH OF HAID 11.	601414000001. [1] ; contains element MER2	Post 1 NIH MGC_16 Homo septems CDNA clara INA CE 22550.2	CO1111970F1 NIH_MGC_16 Home septems CONA clane MACE ASSESSED	throsenous NIH MGC_8 Hamp explens cDNA clone IMAGE:3345480 F	P53620 COATONIED Co14 Homo seplens cDNA clone IMAGE:3035607 3" LET	Homo sablens m.R.N.A. 6. 1214	ws90h03 x1 NC CoAB Co	STRIATIN	Homo saplene raka into CA3816 O43816 O43816	Homo saplens ratis Interest.	Homo septems hardhold and proceed a market of the control of the c	Homo sapiens NF2 none	Homo saplens NF2 next	Homo sapiens culin 4A (CUI 4A) mBNA	SDO each transition of the state of the stat	Homo saplens pleiotrophin (heparin binding growth factor 8, neurite growth manner.	COTO SERVINE LIAL COLOR 17 (P.T.N.) MRNA	60147407251 NIJ 1105	601474072F1 NIH MOC 6211 TITLE BEDIENS CONA CIONE IMAGE:3877121 6	185-01 VAIN COLOR Homo septems cDNA clone IMAGE:3877121 F	EST372729 MAGE resemble 11 Home septems cDNA clone IMAGE:2952384 31	zk45h09.r1 Soares_pregnant ulerus NbHP/I Home saplens cDNA	PIR:S52437 S52437 CDP-discy/diyerd synthase - fruit fiv.	KASCUA.YI NCI_CGAP_Bm52 Homo sepiens cDNA clone IMAGE:2291626 6
Exon Probes	Top Hit Database	Source		Į.	٤	EST HUMAN		EST HUMAN		EST_HUMAN	EST HUMAN	T	Т	Т	EST_HUMAN	Т	Г	EST_HUMAN								$\int$		EST HUMAN G	Т	Г		Г	7	NAMIOT 102
98	salon		I	1	T			7	_		_	ľ	T	$\dagger$	m	Z	-	ŭ	Z	¥	2 6	Z	튁	Z	- 5		N N	ES	ES.	ESI	ES	-	3 8	
	Top Hit Acession		Al 489900 0	7 - 1000 M	A-631820.1	Al732858.1		N99520.1	•	AA011080.1	BE256829.1	BE256829.1	BE268042.1		4W779812.1	4B037768.1		W004608.1	4.0E-61 AF263306.1	NF263306.1	8923209 NT	18000.1	16000.1	3.0E-81 AF077188.1	4508280 VIT	NAME OF THE PARTY	4506280 NT	2.0E-81 BE784636.1	784636.1	V611542.1	7860658.1	1.0E-81 AA040370 4	1.0E-81 BE047898.1	
Most Clarif	(Top) Hit BLAST E	Value	OF P	1 5 E		1.0E-80	20 00	3					5.0E-81	_		10.0		5	2	4.01.61 A 10.10	10 10	3.0E-61 ₹	10.00	<u> </u>	ě	-	φ		20E-81 BE7	<u>ا</u>		81 AA	<u>8</u>	
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	Expression Signet		1.79	1.63		227	0.85			CA:O	3 3	0.63	5.16	•	2.0	8	. 0	2000	100	000	15.97	12.27	14.	-	6.68		20.08	318	3 8	3	8	2.11	10.97	
	ORF SEQ ID NO:			10842		T	14642		12230	14244	14245	13069	1423	11870	13131	-	13558	14019	14020	14251	11285	11286	12405		12957	1205a	12804	12805	13693	13581	3	14354	14479	,
Į.	SEQ ID	2009	3	5812	900		9829		7123	9265	9255	7134	1	8778	8113	$\vdash$	8551	9032	9032	9261	6246	6246	7285	-	E E	7941	7780	7780	0698	8575	-	8375	8200	
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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Hamo septens HSPC288 mRNA, partial cds	Homo sapiene HSPC288 mRNA, partial cds	Human CRFB4 gene, partial cds	Human CRFB4 gene, partial cds	Human CRFB4 gene, partial cds	Homo sapiens mRNA for KIAA1327 protein, partial cds	Homo saplens glutathione percodesse 5 (epididymal androgen-related protein) (GPX5), transcript varient 2, mRNA	Homo sepiens hypothetical protein FLJ20461 (FLJ20461), mRNA	601458531F1 NIH_MGC_66 Hamo sapiens aDNA clane IMAGE:3862086 5	AU144050 HEMBA1 Homo sepiens cDNA clone HEMBA10007523*	Homo sapiens alpha-tubulin Isoform 1 mRNA, complete cds	Homo sapiens amyloid beta (A4) precursor protein (protesse nextr-II, Alzheimer disease) (APP), mRNA	RC2-BN0120-010400-013-f02 BN0120 Homo septems cDNA	Homo sapiens transforming growth factor beta-activated kinase-binding protein 1 (TAB1), mRNA	Homo saplens amyloid beta (A4) precursor protein (protesse nextn-II, Alzheimer disease) (APP), mRNA	al23e05.s1 Soares_testis_NHT Homo sapiens cDNA clone 1343848 3"	RC6-PT0001-190100-021-B02 PT0001 Homo seplens cDNA	Homo sapiens chromosome 21 segment HS21C085	RC1-BN0005-260700-018-g04 BN0005 Homo sapiens cDNA	Homo sapiens neurotrophic tyrosine kinase, receptor, type 2 (NTRK2) mRNA	Homo sapiens mRNA for KIAA0999 protein, partial cds	Homo saplens mRNA for KIAA0999 protein, partial cds	DKFZp434M117_r1 434 (synonym: htes3) Hamo saptens aDNA dane DKFZp434M117 6	Homo sapiens DNA for amyloid precursor probein, complete cds	Homo sapiens glutamate receptor, ionotropio, kainate 1 (GRIK1) mRNA	Homo sepiens mRNA for KIAA1096 protein, partial cds	Homo sapiens mRNA for KIAA1098 protein, partial cds	Homo sapiens wbscr1 (WBSCR1) and wbscr5 (WBSCR5) genes, complete cds, alternatively spiloed and repiloation factor C subunit 2 (RFC2) gene, complete cds	Homo sapiens tumor necrosis factor receptor superfamily, member 5 (TNFRSF5) mRNA	Homo sapiens tumor necrosis factor receptor superfamily, member 6 (TNFRSF5) mRNA
Top Hit Database Source	NT	Z	N	N	IN	IN	_ <del>_</del> =	7	EST HUMAN	EST_HUMAN	TN	_ <del>_</del>	EST_HUMAN	<del>ر</del> ا	_ <u></u>	EST HUMAN	EST_HUMAN	Þ	EST_HUMAN	¥	NT.	Į.	EST HUMAN	F	TN	NT	NT	_F	NT	Į.
rop Hit Acession No.		8.0E-82 AF161406.1				3.1	6715801 NT	8923432 NT		7.0E-82 AU144050.1		4502166 NT	3.0E-82 BE005705.1	5174702 NT	2168			L163285.2	E813232.1	3811			-	J87675.1	4504116 NT	AB029019.1	4B029019.1	AF045555.1	07580	4507580 NT
Most Similar (Top) Hit BLAST E Value	8.0E-82 AI	8.0E-82	8.0E-82 U08988.1	8.0E-82 U08988.1	8.0E-82 U08988.1	8.0E-82	8.0E-82	8.0E-82	7.0E-82 B	7.0E-82	4.0E-82	3.0E-82	3.0E-82	3.0E-82	3.0E-82	3.0E-82	3.0E-82	3.0E-82			2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	2.0E-82	Ш
Expression	4.69	5.26	2.65	. 2.38	1.83	1.39	1.23	0.84	1.04	1.55	100.34	16.31	3.55	6.3	8.88	80.59	96.0	2.02	1.66	2.15	1.39	1.39	1.78	0.85	0.83	1.06	1.06	2.77		1.58
ORF SEQ ID NO:	10077	12001	10330	10855		11515	11681	1	١	12785		10347			10919				11937				11715	13758	14092	14403			L	Ц
Exan SEQ ID NO:	6093	2609	5320	5825	5894	6456	6814	<u> </u>	6421	7651	6299	5334		2800	5878	6054			6849	8214			6843	8759						Ш
Probe SEQ ID NO:	13	107	281	804	876	1459	1617	4121	1424	2693	1632	276	683	779	859	1044	1337	1438	1860	3198	269	265	1647	3756	4112	4426	4426	4720	4908	4908

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	Top Hit Descriptor	•	Description of the second of t	Indus septems melanoma differentiation associated protein-6 (MDA5), mRNA	OUTSTUBBELL NIH, MGC_71 Hamo septens cDNA clane IMAGE:3912207 6	RC4-B10310-110300-016-f10 BT0310 Homo septiens cDNA	Homo sapiens mRNA for KIAA0538 protein, partial cds	10012/3346F1 NIH_MGC_20 Hamp sapiens cDNA clone IMAGE:3814382 F	Ze48f12.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone INAGE-296823.3	QV4-L 10016-271298-068-h11 LT0016 Homo sapiens cDNA no12h01.s1 NCL CGAP_Phe1 Homo sapiens cDNA clone IMAGE:1100497 3' similar to constains At-	Populure sement;  797807x1 NCI_CGAP_Pr28 Home sepiens cDNA clone IMAGE:3847883 3' similar to TR-coverus contacts.	7937807x1 NCI_CGAP_P728 Home saplens cDNA clone MACE-asazone at the control of th	DJ207H1.1;	Human platelet Glycoprotein lib (GPIIb) gene, exons 2-29 hi31hn3 서 Society 하다 구요	SW-YRER HARIN DAXAZA LUNDATI ITTION Septems CDNA clone IMAGE:2833625 3' stmiler to	Homo sapless of the Control of the C	Also apriorite California ZT unknown mRNA	Home series 16ta IIVer splean 1NFLS S1 Home sapiens cDNA clone IMAGE:436080 31	Umaria saprens riypomencal protein FLJ10379 (FLJ10379), mRNA	Hullian euconate denydrogenase fron-protein subunit (schB) gene, exon 6	Floring suprems 26S protessome regulatory subunit (SUG2) mRNA, complete ods	Investigating perfect mapping to chomosome X	Home suprem decoynbonuclease I (DNASE1), mRNA	Homo contons and the CALL MRNA	Homo sapiens mannosidase, beta A, Iyasamal (MANBA) gene, end ubirnitin continuets.	(UBEZD3) genes, complete cds	001511580F1 NIH_MGC_71 Home septens cDNA clone IMAGE:3913195 6	ES179542 Placenta i Homo sapiens cDNA similar to similar to endogenous retrovirus ERV9 np87c97.s1 NCL CGAP Thy Homo sapiens cDNA clare INA CE 443222	spelitive element;	0664905.81 Scares_testis_NHT Homo sapiens cDNA clone IMAGE:1621592 3' similar to TR:092614 C92614 MYELOBLAST KIAA0218.;	
	Top Hit Detabase Source				ESI TOMAN	TOWAN		ES LICHAN	HOMAN	HOMAN		NAME OF THE PERSON OF THE PERS	EST HUMAN		EST HUMAN	Т	EST HIMAN	Т						-   		T	T	Т	EST HUMAN	EST_HUMAN C	
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	Top Hit Acessian No.		11545024 NT	Į,	REDRASBA 4	AB04440.0	8 0F-R3 BE383079 4	8 OF 83 NASOE4 4	AW385520 4	7.0E-83 AA584855 1	7.0E-83 BF224813 4	7 05 02 0500000	A33300 4		6.0E-83 AW573088.1	6.0E-83 AF231919.1	6.0E-83 AA701457.1	11430241 NT	17883 1	5.0E-83 AF006305 4	1133207 2	4885100 NT	4557013 NT	4557013 NT	8	000	A36044 4			492.1	
	Most Similar (Top) Hit BLAST E		1.0E-82	1 0F-82 RE	1.0F-82 REG	1 OF 82 ABO	8 OF R	A OF A2	7.0E-83 AW	7.0E-83	7.0E-83	7 05 02	6.0F-83 M833		6.0E-83	6.0E-83	6.0E-83	6.0E-83	5.0E-83 U17883 1	5.0E-83	5.0E-83 AL 13	5.0E-83	5.0E-83	5.0E-83	4 05 83 4592	4 OF 83 B	3 0E-82 A A38924 4	2 2 2	3.0E-83 AA632654.1	2.0E-83 AA983	
	Expression Signal		1.67	1.57	32	114	4.62	43	1.67	1.62	6.92	1.564	16		7.97	1.02	1.01	0.82	1.85	2.85	1.13	0.99	14.02	14.02	1.78	1.08	3.79	;	4	1.7	
	ORF SEQ ID NO:		10616		11305	11308		11709	11386				10458		11822		13026	13507			13566	13819	14857	14858	10867	13469	-			11841	
	SEQ ID	1				9265	8386	L	6337	7814	9855	10001	5437	-	0/42	7995	8014	<b>8</b>	6949	104	8560	8813	9886	9886	2862	8443	26897	7883		6756	
	SEO ID		280	1189	1288	1267	1389	1642	1339	2794	4670	5036	401	47.60	7	2 2	9887	3483	832	1997	3553	3810	4807	4807	88	3435	885	2705		1784	

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Single Exon Probes Expressed in HBL100 Cells	Top Hit Descriptor	otistagos as Sources_trestis_NHTT Homo septiens cDNA clone (MAGE:14534502 %; e/msitre-te-TE-Cases.	QBZ014 MYELOBLAST KIAA0Z16.;	2848112.81 Soares fetal liver spleen 1NFLS Homo septens cDNA clone IMAGE:286823.3	INCO-E 10046-280600-013-H12 ET0046 Homo saplens cDNA	Homo sapiens sal (Drosophila)-like 1 (SALL1), mRNA	Homo sepiens chromosome 21 segment HS21C002	Homo seplens hematopolatic progenitor cell antigen CD34 precursor (CD34) mRNA, partial col-	Homo seplens ankyrin repeat-containing protein ASB-2 (LOC51676), mRNA	Fromo saplens ankyrin repeat-containing protein ASB-2 (LOCS1676), mRNA Homo saplens hydroxyacyt-Coenzyme A dehydrogenasa/3-ketoscyt-Coenzyme A thiolesa/ennal Coenzyme A	nycreuse (intrinctional protein), beta subunit (HADHB) mRNA Homo saplens hydroxysovi-Costzone A dehydroxysovi-Costzone A	hydratase (trifunctional protain), beta subunit (HADHB) mRNA	601507375F1 NIH_MGC_71 Homo sepiens cDNA clone IMAGE:3908754 F	Rattus nonegicus brain specific cortactin-binding protein CBP90 mRNA nartial col-	H.sapiens gene for mitochondrial dodecency-CoA delta-isomerase, excn 3	Morning and the second and second	Solestens amyond beta (A4) precursor protein (protesse next-II, Alzheimer disease) (APP), mRNA	GE:395863 6'			wed and at Secret for the second or an action of the second control in the second seco	9		done IMAGE:2302086 3' similar to	SWINNIC HUMAN 043847 NARDILYSIN PRECURSOR;	Home sapens powersse (DNA-directed), alpha (70kD) (POLA2), mRNA	House septents myostin light chain kinase Isoform 2 (MLCK) mRNA, complete cats	ds	Novel human mRNA contenting 1 (PCM1) mRNA	2 (DLEC1, ORCTL3, ORCTL4 genes,
Exon Probes	Top Hit Detabase Source		EST HUMAN	EST HUMAN	NAMOR 161				L	5			HOMAN		LZ.		T HIMAN	Т	Т	Т	Т	Т	7	7444	אלאטרי					
Single	Top Hit Acession No.	440094004	_			AI 1832	. I 🔻					4328		4 OE 92 775000 4	T	4502168/NT	7.0E-84 BE901209.1	T	Τ	Τ	Γ	T	5.0E-84 AF109718.1		4505028		3.0E-84 AF026200 1	3855	9898	
	Most Similar (Top) Hit BLAST E Value	2.05.82	2 OF 43	2.0E-83	2.0E-83	20E-83	2.0E-83	2 OF-83	2.0E-83	1.0E-83		1.01-83	4 OF B2	2010	1,05-80	1.0E-83	7.0E-84	8.0E-84	6.0E-84	6.0E-84	6.0E-84	6.0E-84/	5.0E-84 /	4.0F-84 At	4.0E-84	4.0E-84	3.0E-84	3.0E-84	3.0E-84 AL	3.0E-84 A
	Expression Signal	1.7	3.01	2.73	222	0.74	4.16	6.32	6.32	231	200	132	531	3.74		2.36	4.14	2.87	2.87	3.11	0.98	0.8	1.24	271	1.93	1.73	1.77	1.89	3.88	76.0
	ORF SEQ ID NO:	11842					14185	14487	14488	11435	11438	12683		l		14703	13714	11313	11314	12429	14984	10745		11434	14760	14761	10380	12000	12046	13529
	SEQ ID NO:	6756	6873		8211				9507	6384	6384	7648	8780	9116		9718	8711	6274	6274	7308	10015	6728	7364	6383	9776	2225	5369	9069	6944	8518
	Probe SEQ ID NO:	1764	1884	2779	3195	3687	4210	4517	4517	1387	1387	2585	3777	4122		4733	3707	1276	1278	2334	5044	704	2945	1386	4782	4793	314	1920	1958	3610

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Table 4
Single Exon Probes Expressed in I

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Single Exon Probes Expressed in HBL100 Cells	Top Hit Descriptor	Home senters Vinted to	(344 Proget speed 279 Lee Progettis predicts protein (XLRS1) mRNA, complete cds	CAM BIDISE (80400 070 Los Branch)	Home earlies mindle feet of th	Historian Complete and Commence of the Complet	Homo sablens triansectin short forform (TSN) mBNA	Homo sepiens tyrosine 3-monoxygenase/typtophan 5-monoxygenase activation protein, zeta polypeptide	Homo sariens complement comments (Agrants	am85b11 at Stratages enter both 544 U	601308006F1 NIH MGC 44 Home senime chiva class 14.0 Executive	Homo sapiens melcentrioler metaries 4 (DCMA) DMA	my 2808.81 NCI COAP SS1 Home england a Day of Long 114 CE (2000)	Homo saniens 959 to contra between AMI 4 2-1 CPB4	DKFZ0434N0323 rt 434 (sunovambles) Lorre collection CNR 21 CZ segment 13	DKFZp434N0323 r1 434 (swowyn hies3) Homo septems cDNA close DK-Zp434N0323 6	Hamo septens 959 to contra between AMI 1 and CRD1 on changes 24 222	RC4-BT0311-141299-012-008 BT0311 Homo senions conta	RC4-BT0311-141299-012-008 BT0311 Homo saniems CONA	Homo saplens chromosome 21 segment HS21Chho	Homo sapiens nuclear protein Skip mRNA, complete cds	Homo sapiens nuclear protein Skip mRNA, complete cds	Human plasminogen gene, expn 7	Human plasminogen gene, exon 7	Homo sapiens DKFZp434P211 protein (DKFZP434P211), mRNA	Homo saplens nucleolar GTP ase (HUMAUANTIG), mRNA	Homo sapiens chromosome 21 segment HS21C080	Human ornithine decarboxylase gene, complete cds	Human omithine decarboxylase gene, complete cds	Homo sapiens chromosome 21 segment HS21C068	Homo sapiens ribosomai protein L27 mRNA, complete cds	Homo sapiens chromosome 21 segment HS21C084	Homo sapiens protein phosphatase 2A BR gamma subunit gene, exon 6	
xon Probes	Top Hit Detabase Source		EST HIMAN	EST HIMAN	LN	F	Z		: 5	EST HUMAN	EST HUMAN		EST HUMAN	IN	EST HUMAN	EST HUMAN	Z	EST HUMAN	EST HUMAN	NT	NT	NT	NT	N	E	ΙT	NT	NT	TN	NT	LN		П	
a elibuic	Top Hit Acession No.	-014459.1	T	2.0E-84 BE695397.1	T		1.0E-84 AF114488.1	36.	11427831 NT	984379.1	392137.1	7197	720851.1	Γ	1.0E-84 AL043314.2	Γ	1.0E-84 AJ229041.1 N						3282.1	3282.1	7657020 NT		163280.2			.2			3.0E-85 AF096157.1 N	
	Most Similar (Top) Hit BLAST E Veitue	3.0E-84 A	2.0E-84	2.0E-84	2.0E-84	2.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84	1.0E-84 BE	1.0E-84	1.0E-84 AA	1.0E-84/	1.0E-84/	1.0E-84 /	1.0E-84	1.0E-84 /	1.0E-84	9.0E-85	9.0E-85 U51432.1	9.0E-85 U51432.1	9.0E-85 M33282.1	9.0E-85 M3	9.0E-85	9.0E-85	9.0E-85 AL	9.0E-85 M	9.0E-85 M33764.1	9.0E-85 AL	7.0E-85 L05094.1	5.0E-85 AL	3.0E-85	100 100
	Expression Signal	5.61	4.68	4.68	8.83	1.41	1.21	16.96	123	2.78	1.8	1.8	2.55	4.33	2.93	2.93	2.67	0.71	0.71	2.75	10.8	10.8	=	+	2.05	0.91	1.08	1.44	1.44	1:1	34.29	2.11	0.77	000
	ORF SEQ ID NO:	13873	12141	12142	L	12932	10376	10583		11311	12088	12258	13675	14271	14534	14535	14271	14907	14908		11094	11095	11598	11599	11702	13717	14109	14615	14616	14726	11155	12364	11320	44040
	ш W ~		7029		7892		6365	5579	5734	8272	6984	744	8670	9284	8549	9549	9284	8929	8929	2968	6065	6065	6241	L#C0	9632	8715	9125	9624	9624	9741	6125	7247	6279	0757
	Probe SEQ ID NO:	3662	2047	2047	2873	2892	310	645	710	1274	2001	2162	3865	4292	4561	4561	4756	4952	4952	827	1056	1056	1043	2 5	1635	3711	8 8	4639	4639	4757	1119	227	1280	4772

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	801189704F2 NIH MGC 7 Homo sepiens CDNA clone IMAGE:3633816 6	House services E has conto monthly 24 (FBXO24) mRNA	ndili sepesis i tur din promi e 10 menu	Hamo septems 1-box only process 2+ (1-bx/2.2-), service 3 (OB12072) mRNA	Homo sapiens offactory receptor, Tamily 12, sumanilly D, Intelliber 2 (Ch. 1222), Inc.	Homo sapiens KIAA0929 protein Msx2 Interacting nuclear target (MINT) homolog (KIAA0929), mRNA	Homo sapiens intersectin 2 (SH3D1B) mRNA, complete cds	Homo saplens CGI-201 protein (LOC51340), mRNA	Homo sapiens apolipoprotein C-II (APOC2) minuta	Homo sapients apolipoprocein Chi (NFOCZ) illusos	Human DNA polymerase being gene, aware 12 and 13	Homo sapiens similar to rat tribaja a menuniane grychycem.	Human Ku (propeu) supumit mitures, continues cas	Homo septens plasmingen (TLC) mirans	Homo sepiens chromosome zi sepiment nozi over	601591416F1 NIH_MIGC_/ India 8g/fais cDNA clone IMAGE:3866021 6	0014626 I/T NIT 1910 CV There explans CONA done IMAGE:3866021 5	601462817F1 NIH MCC 20 Home septems CDNA clone IMAGE:2967690 5	See 1 See 1 Acres Parethyrid fumor NbHPA Homo sapiens cDNA clone IMAGE: 1403559 3'	glocino, st. 3 cares, parathyroid fumor NDHPA Homo sapiens cDNA clone IMAGE: 1403569 3*	Homo seniens avodutarate dehydrogenase (lipoemide) (OGDH) mRNA	Homo sapiens 24 kDs intrinsic membrane protein (PMP24), mRNA	601072594F1 NIH MGC 12 Homo septens cDNA clone IMAGE:3458830 6	EST177232 Jurkat T-cells VI Homo saplens cDNA 6" end	Hamo saplens chramosome 21 segment HS21C003	VZ19808.r1 Sogres_multiple_sclerosis_ZNbHMSP Homo sepiens cONA clone IMAGE:263478 o	Human endocenous retrovirus, complets genome	Home saplens mRNA for KIAA1277 protein, pertial cds	FST378215 MAGE resequences, MAGI Homo saplens cDNA	Home seriens hisophosphatidic acid acyltransferase-delta (LPAAT-delta) mRNA, complete cds	Homo saciens hysophosphatidic acid acyltransferase-delta (LPAT-delta) mRNA, complete cds	MARZOB X1 NCI CGAP GC6 Homo septens cDNA clone IMAGE:2916542 3'	Homo sapiens cAMP-specific phosphodiesterase 8A (PDE8A) mRNA, partial cds	
	Top Hit Detabase Source	COT LIBIAN	Name I I		E	F		N	11	<del> </del>	누	¥	Ę	Ä	Ę	Ę	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	ESI HUMAN		CCT UI MAN	EST HIMAN	TN	EST HIMAN	TIV	z	COT LIMAN		į į	N. COT LIMAN	NT TOWN	
	Top Hit Acession No.	7,0072	1.00	11024695INI	11024695 NT	7363442 NT	7857268 NT	8540.1	7706205 NT	5174775 NT	3174775	J10525.1	7857468 NT		4505880	63284.2	94306.1	18392.1	318392.1	274217.1	7.0E-86 AA860801.1	AA86080		IN SESCONS	A A 2002 4 4	2.0E-60 AASUGEOT: 1	AL 100203.4	2.0E-86 N369/7.1		2.0E-86 AB033103.1	2.0E-86 AW906142.1	2.0E-86 AF156776.1	2.0E-86 AF156776.1	2.0E-86 AW516742.1	Aronago.
	Most Similar (Top) Hit BLAST E Veiue	1000	3.0E-80 BE.20	3.0E-85	3.0E-85	3.0E-85	2.05.85	2.0E-85 AF24	2.0E-85	2.0E-85	2.0E-85	2.0E-85 U10525.1	2.0E-85	2.0E-85 M30838.1	2.0E-85	2.0E-85 AL1	1.0E-85 BE?	1.0E-85 BE	1.0E-85 BE6	9.0E-86 BE		7.0E-86													2.0E-80/AL
	Expression Signal		1,28	1.48	1.48	-	07.0	1 98	1.33	7.62	7.62	4.1	10.24	2.18	5.78	76.0	3.19	5.67	5.67	54.78															3.42
	ORF SEQ ID NO:		14164	14714	14715	14775	8000	10880	11425		11448	12265		12890			L	12423			10972		11312			3 10332					13359	13671	13672	Ш	14825
	Exam SEQ ID NO:		98	9728	9772	9783	1	2504	8377	6392	6392	7148	6315	7975	9200	9740	7200		L	L				8972					3 7108	7184	1 8341	1 8666	1 8666		5 9630
	Probe SEQ (D NO:		4188	4743	12/43	4809	1	3 3	280	1385	1395	2169	2750	2956	4207	4755	2223	2329	2329	1404	923	923	1275	5001	211	264	411	1170	2128	2207	3331	3661	3661	3927	4645

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo saplens NADH dehydrogenase (ubiquinone) Fe-S protein 1 (76tD) (NADH-coenzyme Q reductase) (NDUFS1) mRNA	Homo saplens (ibulin 6 (FBLN6) mRNA	Human gamme-dutamy transpeditions mRNA, complete ods	Homo sapiens chromosome 21 segment HS21000s	Homo sapiens chromosome 21 segment HS21C009	Homo sapiens hypothetical protein (LOC51318), mRNA	Homo sapiens hypothetical protein (LOC51318), mRNA	Homo saplens chromosome 21 segment HS21C100	O.cuniculus mRNA for elongation factor 1 alpha	7h85f02.x1 NCI_CGAP_Co16 Homo septens cDNA clone IMAGE:3322779.3'	7h85f02x1 NC  CGAP Co18 Hamp septens cDNA clans IMAGE:3322776 3	Homo sapiens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA	EST96094 Testis I Homo saplens cDNA 5' and	Homo sapiens chromosome 21 segment HS21C010	Homo sapiens mRNA for KIAA1414 protein, partial cds	Homo saplens mRNA for KIAA0456 protein, partial cds	Homo sapiens CGI-60 protein (LOC51626), mRNA	Homo saplens CGI-60 protein (LOC51626), mRNA	Homo sapiens myekold/lymphtoid or mbed-lineage leukemia (trithorax (Drosophila) homolog); translocated to, 4 (MLLT4) mRNA	Homo sapiens high-mobility group (nonhistone chromosome) protein 4 (HMG4) mRNA	QV0-BN0148-050600-254-e03 BN0148 Hamo sepiens cDNA	AU116935 HEMBA1 Homo sapiens cDNA clone HEMBA1000307 6	CM0-TN0038-150900-552-h08 TN0038 Homo sepiens cDNA	Homo saplens putative glycolipid transfer protein (LOC51054), mRNA	PM2-CT0265-141099-001-904 CT0265 Home sepiens cDNA	PM2-CT0265-141099-001-g04 CT0285 Homo saplens cDNA	Human mRNA for T-cell cyclophilin	Homo sapiens neureskin III (NRXN3) mRNA	Homo sapiens protesse Inhibitor 4 (kallistatin) (PI4) mRNA	Homo saplens double stranded RNA activated protein kinase (PKR) gene, exon 12	Homo sapiens mRNA for KIAA1389 protein, partial cds	Homo sapiens mRNA for KIAA1399 protein, partial cds
	Top Hitt Database Source	_ <b>5</b>	¥	¥	N <sub>T</sub>	Į.	N P	F	F	N	EST_HUMAN	EST HUMAN		EST HUMAN	E	Z	Ę	F	NT.		Z	EST_HUMAN	EST_HUMAN	EST_HUMAN	IN	EST_HUMAN	EST_HUMAN	N.	Į.	NT	L	N	7
	Top Hit Acession No.	4828855 NT	5453649 NT	20492.1	1.0E-86 AL163209.2		7706181 NT	7706161 NT	1.0E-86 AL163300.2		BF063211.1	F063211.1	7657213	5.0E-87 AA382811.1			AB007925.1	7706299 NT	7706299 NT	5174574 NT	4885420 NT		2.0E-87 AU116935.1	F376311.1	7705883 NT	1.0E-87 AW361977.1	7.1		4758827 NT	5453887 NT	F167485.1		9.0E-88 AB037820.1
	Most Similar (Top) Hit BLAST E Vatue	1.0E-88	1.0E-86	1.0E-86	1.0E-86	1.0E-86	1.0E-86	1.0E-86	1.0E-86/	8.0E-87 X82245.1	7.0E-87	7.0E-87 B	6.0E-87	5.0E-87	4.0E-87	4.0E-87	4.0E-87	4.0E-87	4.0E-87	4.0E-87	2.0E-87	2.0E-87 BI	2.0E-87	2.0E-87 B	1.0E-87	1.0E-87	1.0E-87	1.0E-87 Y00052.1	1.0E-87	9.0E-88	9.0E-88 AI	9.0E-88 /	9.0E-88/
	Expression Signal	2.08	1.45	2.68	1.17	1.17	1.22	1.22	6.37	131.14	1.99	1.99	0.78	2.38	1.12	18.02	1.85	1.2	1.2	1.88	6.54	1.17	0.88	99.0	1.89	1.69	1.69	13.27	2.3	0.7	6.3	2.48	2.48
	ORF SEQ ID NO:	11621		13202	13258	13259	13852	13853	14121			12326	13482	11176	11000	11190	12071	12444	12445	13414	12773		13703	14727		11460	11461	13640	13661	10961	11125	11377	11378
	SEQ ID	6659	8105	8180			8844	8844	9137	5512	7210	7210	8456	6145	5967	6157	6962	7329	7329	8391	7661	7897	8700	9742	7698	6403	6403	8635	8655	5926	9609	6328	6329
	Probe SEQ ID NO:	1562	3089	3164	3222	3222	3842	3842	4142	476	2233	2233	3448	1140	951	1153	1980	2355	2355	3383	2704	2878	3696	4758	1163	1406	1406	3628	3649	608	1089	1331	1331

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Homo sepiens chromosome 21 segment HS21Co09	H.saplens ECE-1 gene (exen 9)	H. saplens ECE-1 gene (excn 0)	Homo sapiens KIAA0083 gene product (KIAA0083), mRNA	K9719F Human fetal heart, Lumbda ZAP Express Homo sapiens cDNA clone K9719 5' similar to ZINC FINGER PROTEIN HZF1	Homo saplens intersectin short tectorm (ITSN) mRNA, complete cds	Homo saplens intersectin short isoform (ITSN) mRNA, complete cds	Homo sapiens intersectin short isoform (ITSN) mRNA, complete cds	wd68h08.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:2336769 3' similar to contains Alu repetitive element, contains element MER22 MER22 recettive element:	Homo sapiens Intersectin short isoform (ITSN) mRNA, complete cds	PM1-TN0028-050900-004-f10 TN0028 Homo saplens cDNA	PM1-TN0028-050900-004-f10 TN0028 Homo sapiens cDNA	Homo sapiens hypothetical protein FLJ21634 (FLJ21634), mRNA	Homo saplens zinc finger protein 259 (ZNF259) mRNA	za48f12.s1 Soares fetal liver spleen 1NFLS Home sepiens cDNA clone IMAGE:285823 3'	Homo saplens a disintegrin and metalloproteinase domain 23 (ADAM23) mRNA	Homo saplens a disintegrin and metalloproteinase domain 23 (ADAM23) mRNA	Homo sapiens hypothetical protein FLJ20220 (FLJ20220), mRNA	Homo saplens Calsenilin, presenilin-binding protein, EF hand transcription factor (CSEN), mRNA	Homo saplens SNARE protein kinase SNAK mRNA, complete cds	Homo sapiens SNARE protein kinase SNAK mRNA, complete cds	Homo sapiens SNARE protein kinase SNAK mRNA, complete cds	Homo sapiens dynein, exchemal, light polypeptide 4 (DNAL4), mRNA	601142409F1 NIH_MGC_14 Hamo sapiens cDNA clone IMAGE:3506188 67	Homo sapkens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA	Homo sapiens hormonally upregulated neu tumor-associated kinase (HUNK), mRNA	Homo saplens complement component 8, beta polypeptide (C8B) mRNA	DKFZp434E248_r1 434 (synonym: htes3) Hamo sapiens cDNA clone DKFZp434E248 5	Homo sapiens inner membrane protein, mitochondrial (mitofilin) (IMMT), mRNA	Homo sapiens serine/threonine-protein kinase PRP4 homolog (PRP4) mRNA	Homo sepiens ubkquitin-conjugating enzyme E2L 3 (UBE2L3) mRNA	Homo sapiens ubiquitin-conjugating enzyme E2L 3 (UBE2L3) mRNA
Top Hit Datebase Source	E	F	N T	ラ	ST HUMAN	N.	Z	Z	ST HUMAN	F	EST_HUMAN	EST_HUMAN	F	77	EST_HUMAN	Ę		7	<b>1</b>	NT	L/L	N	<b>-</b>	EST HUMAN	レフ	L)	77	EST_HUMAN	イ	トフ	5	F
Top Hit Acession No.	1163209.2		9.0E-88 X91929.1	7661887 NT	5.0E-88 N89399.1	  =		6.0E-88 AF114488.1					11545800 NT	4508020 NT	3.0E-88 N66951.1	4501912 NT	. 4501912NT	11429300 NT	7305198	-246219.1		2.0E-88 AF246219.1	1666	8.0E-89 BE311557.1	7657213 NT	7657213 NT		L045748.1	5803114 NT	4508124 NT	4507788 NT	4507788 NT
Most Similar (Top) Hit BLAST E Value	9.0E-88 A	9.0E-88	9.0E-88	5.0E-88	5.05-88	5.0E-88	6.0E-88	5.0E-88	5.0E-88 AI	5.0E-88	4.0E-88	4.0E-88	3.0E-88	3.0E-88	3.0E-88	3.0E-88	3.0E-88	3.0E-88	2.0E-88	2.0E-88 AI				8.0E-89	7.0E-89	7.0E-89	7.0E-89	7.0E-89 A	6.0E-89	6.0E-89	6.0E-89	6.0E-89
Expression Signal	0.89	3.04	3.04	1.98	4.62	0.71	0.76	0.76	2.58	0.7	1.13	1.13	1.4	2.28	4.85	0.69	0.69	3.64	1.27	1.92	4.8	0.99	2.35	1.49	1.24	1.24	3.49	5.7	2.12	1.98	5.05	5.05
ORF SEQ ID NO:	13557	14126	14127		12650	12967	12979	12980		13493	11355	11356	10764		12920	14096	14097				١	13415			10486	10487	14700	14746	11047			12457
Exon SEQ ID NO:	8549	9142	9142	6783	7531	7950	7960	7960	8316	8466	6307	6307	5745	8929	7896	9112	9112	9342	6029	6582	9029	8392	9293	7620	5468	5468	9715	9758	6017	7130	340	7340
Probe SEQ ID NO:	3542	4147	4147	1792	2568	2931	2941	2941	3305	3458	1309	1309	722	1778	2877	4118	4118	4351	1019	1585	171	3384	4301	2661	430	430	4730	4774	1007	2151	2366	2366

Page 146 of 209 Table 4 Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	•	Homo saplens HSPC169 protein (HSPC169), mRNA	Homo saplens HSPC019 protein (HSPC019), mRNA	Homo saplens mRNA for KIAA0408 protein, partial cds	Homo sapiens mRNA for KIAA0408 protein, partial ods	TCBAP2E0383 Pediatric pre-B cell acute lymphoblastic feukemia Baykor-HGSC project=TCBA Homo sapiens	TCBAP2E0383 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo saplens	EST388290 MACE resentations MACN Home sections COMA	Homo saplens PXR2b protein (PXR2b) mRNA	Homo saplens PXR2b protein (PXR2b) mRNA	Homo sapiens mRNA for KIAA1342 protein, partial cds	998008.x1 Soures, NR_T_GBC_S1 Homo septens cDNA clone IMAGE:1843022.3' stimiter to gb:J04131 GAMMA-GLUTAMYLTRANSPEPTIDASE 1 PRECURSOR (HUMAN) contents a fut repositive advanced.	Homo saplens topolsomerase-related function protein (TRF4-2) mRNA, partial cds.	H. sapiens HCK gene for tyrosine kinase (PTK), exons 10-11	H.saplens HCK gene for tyrosine kinase (PTK), exons 10-11	Hamo sapiens chromosome 21 segment HS21C003	Homo sapiens GGT gene, exon 5	Homo sapiens chromosome 21 segment HS21C048	Homo sapiens chromosome 21 segment HS21C046	7e36f08.x1 NCI_CGAP_Lu24 Home septens cDNA clone IMAGE:3284583 3'	7636108.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3284563 3'	Homo sepiens calcium channel alpha1E subunit (CACINA1E) gene, exons 7-49, and partial cds, atternatively spliced	H.sapiens ECE-1 gene (exon 6)	H.sapiens ECE-1 gene (exon 6)	Homo saplens hypothetical protein FLJ10388 (FLJ10388), mRNA	Homo saplens hypothetical protein FLJ10388 (FLJ10388), mRNA	Homo saplens TCL6 gene, exon 1-10b	Human gamma-aminobutyric acid transaminase mRNA, partial cds
Top Hit Databese		¥	¥	N	N.	FST HIMAN	EST LIBIAN	EST HUMAN	L	Z	F	EST HUMAN	¥	Ę	IN	LN L	TN	TN	ĪN	EST_HUMAN	EST_HUMAN	NT N	ĘN	IN	FZ	TN	NT	NT
Top Hit Acession No.		7881817 NT	7881737 NT	AB007866.2	AB007866.2	BE244323 1	F244328 4	AW978181.1	7706870	T706670	AB037763.1	1222095.1	F089897.1	(58742.1		L163203.2	J007378.1		8.0E-90 AL163246.2	E670561.1	E670561.1	7.0E-90 AF223391.1	91926.1	01926.1	TN 8922398	8922398	B035344.1	J80226.1
Most Similar (Top) Hit BLAST E	Value	6.0E-89	6.0E-89	6.0E-89	6.0E-89	5.0E-89	8.0F.80	3.0E-89	2.0E-89	2.0E-89	2.0E-89	. 2.0E-89 A	2.0E-89 A	2.0E-89 X	2.0E-89 >	2.0E-89	2.0E-89	8.0E-90	8.0E-90	8.0E-90 B	8.0E-90 B	7.05-90	6.0E-80	6.0E-90	9.0E-90	6.0E-90	5.0E-90 A	5.0E-90
Expression Signal		0.75	0.91	3.9	3.9	3.31	331	0.95	0.87	78.0	0.68	1.44	1.18	4.76	4.76	1.09	1.05	3.19	2.72	3.66	3.66	4.22	1.08	1.08	9.58	9.58	78.69	2.39
ORF SEQ ID NO:			14324	14470	14471	14852	14853		10462		10564	12847	14005	14014	14015	14198		11084		11357	11358		13024	13025	14085	14088		11210
SEQ ID					9493	9882	9882		5442	5442	5561	7831	8018	9026		9217				7744	7744	5846	8012	8012	6606	8089	5220	6178
Probe SEQ ID NO:		3446	4354	4503	4503	4903	4903	2807	127	127	526	2811	4022	4030	4030	4223	4367	1046	1047	1310	1310	826	2994	2894	4105	4105	154	1173

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	998608x1 Soares NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE:1843022 3' similar to gb:J04131 GAMMA-GLUTAMYLTRANSPEPTIDASE 1 PRECURSOR (HUMAN);conteins Alu repetitive element;	4996008.x1 Soares_NFL_T_GBC_S1 Homo sepiens cDNA clone IMAGE:1843022 3' similar to gb.J04131 GAMMA-GLUTAMYLTRANSPEPTIDASE 1 PRECURSOR (HUMAN):conteins Alti mandillus administ	Homo sapiene intersectin long isoform (ITSN) mRNA, complete cds	Homo saplens chromosome 21 unknown mRNA	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens myosin phosphatase, target subunit 1 (MYPT1), mRNA	H. sapiens gene encoding discodin receptor tyrosine kinase, exon 18	Homo sapiens DNA for emyloid precursor protein, complete cds	Homo sapiens mRNA for KIAA1244 protein, partial cds	Human prohormone converting enzyme (NEC2) gene, exon 8	Homo sapiens collegen, type XII, alpha 1 (COL12A1), mRNA	601087378F1 NIH_MGC_10 Homo sapiens cDNA clone IMAGE:3453834 5	Homo sapiens high-mobility group (nonhistane chromosomal) protein 17 (HMG17), mRNA	Homo sapiens high-mobility group (nonhistone chromosomal) protein 17 (HMG17), mRNA	qc54c02.x1 Soeres_placenta_8tx5weeks_2NbHP8tx9W Homo sapiens cDNA clone IMAGE:1713410 3* similar to SW:CLF3 MOUSE P23275 OLFACTORY RECEPTOR OR3	Homo sapiens mRNA for KIAA0289 gene, partial cds	Homo sapiens GRB2-related adaptor protein (GRAP) mRNA	Homo sapiens emyloid beta (A4) precursor protein (protesse nextn-fl. Alzheimer disesse) (APP). mRNA	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens mRNA for T-box transcription factor (TBX20 gene), partial	Homo sapiens mRNA for T-box transcription factor (TBX20 gene), partial	Homo sapiens ALR-like protein mRNA, partial cds	Homo saplens ALR-like protein mRNA, partial cds	Homo sapiens Kruppel-like factor 7 (ublquitous) (KLF7), mRNA	Homo sapiens protein phosphatase 2A BR gamma subunit gene, exon 3		601159563F2 NIH_MGC_63 Hamp septens cDNA clane IMAGE:3511118 6"
± 8	Database Source	EST HUMAN	EST HUMAN	Т	Z	Ę		NT		TN.	L		EST_HUMAN			EST HUMAN	Т			NT	N	N.	N N		NT				EST_HUMAN
	Top Hit Acession No.	A1222086.1	A1222095.1 E	AF114487.1	AF231920.1	AF231920.1	4505316 NT	CB9033.1	D87675.1	AB033070.1	M95967.1	TN 7779	BE537913.1 E	5031748 NT	5031748 NT	Al138213.1 E	AB006627.1	5729855 NT	4502168 NT	AF231920.1		AJ237589.1	AJ237589.1		AF264750.1	7828			BE379884.1
Most Similar	(Top) Hit BLAST E Vatue	5.0E-90	6.0E-80	5.0E-90	4.0E-90	4.0E-90	4.0E-90	4.0E-90 )	4.0E-90	4.0E-90	4.0E-90	4.0E-90	2.0E-90	2.0E-90	2.0E-90	2.0E-90	2.0E-90	2.0E-90	1.0E-90	1.0E-90	1.05-90	1.05-90	1.0E-90	1.05-90	1.0E-90	1.0E-90	1.0E-90	1.0E-90	1.05.90
	Signet	1.48	1.48	1.82	2.4	2.4	3.28	9.18	4.85	1.95	1.82	0.92	4.2	71.49	71.49	1.88	76.0	8.45	5.3	1.98	1.9	1.73	1.73	11.11	11.11	3	2.29	2.29	4.67
100	OKI- SEQ ID NO:	11867	11868	12588	10369	10370					14641	14958	10290	11191	11192	13757	14524	14724	10348	10436	10436	10723			10762			11330	
Exon	SEQ ID NO:	8778	6778	7452	5357	5357	8078	6646	9512	9638	9658	9981	5276	6158	6158	8758	9538	8238	5333	7693	7693	5710	5710	5743	5743	6100	6286	6286	6625
Probe	SEQ ID	1784	1784	2484	300	300	1070	1650	4522	4653	4673	5010	213	1154	1154	3755	4550	4754	274	373	374	989	989	720	720	1083	1287	1287	1628

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo espiens similar to SALL1 (sal (Drosophila)-like (LOC57167), mRNA	Home earliers chromosome 8 doen reading frame 2 (C8ORF2), mRNA	Umms emissione mRNA for KIAAAARR protein partiel ods	TOURS SEPTION TO THE	Homo sepiens mixing to ruly a second partial control (ii 4RAP) cene, each 8, effertiative excits 9	Homo sapiens solution interisularity to agricultural accessory process, (***).	HINADORS 381 I Ner HebG2 call line. Homo septems cDNA done s3813'	Social Spares fotal liver spleen 1NFLS S1 Homo sapiens cDNA clane IMAGE:448015 3	DBUILD-13 COM AS LINES CANING COME Y TRANSCORT 5	AU1435259 1787A1 Home emittee of NA Jame 778AA1002087 5	AU143538 1 187A1 India espisis control control (C220RE5) mRNA	Homo sapiens diffundaments and the following from 5 (C220RFS) mRNA	Homo sapiens chromosome 22 great teating traine of O22011 01, 11, 11, 11, 11, 11, 11, 11, 11,	Homo sepiens lysophosphaudic acid acyluansishassociata (L. M.).	Homo sapiens lysophosphaudre acid acyluanoleaso com (2.1017)	Homo sapiens source carrier raining 1, carrent contraction of the carrier of the	Home sapiens source carrier taining 4, anion contracts, more and the sapiens of sapiens and sapiens are sapiens and sapiens and sapiens are sapiens and sapiens and sapiens are sapiens and sapiens and sapiens are sapiens and sapiens and sapiens are sapiens and sapiens and sapiens are sapiens and sapiens and sapiens are sapiens and sapiens are sapiens and sapiens are sapiens and sapiens are sapiens and sapiens are sapiens and sapiens are sapiens and sapiens are sapiens and sapiens are sapiens and sapiens are sapiens and sapiens are sapiens and sapiens are sapiens and sapiens are sapiens and sapiens are sapiens and sapiens are sapiens and sapiens are sapiens are sapiens and sapiens are sapiens are sapiens are sapiens are sapiens and sapiens are sapiens are sapiens are sapiens are sapiens are sapiens are sapiens ar	Homo sapiens unquiui rougugating bii voormat Hoodonsa	Home sapiens circuitisating 21 against 1921.	India equals in which the state of the state	Home suplems minuted to the fact of the protein many, complete cds	Luman K. (AZOLABA) submit mRNA, complete cds	Home earliers, chromosome 21 segment HS210085	Homo saviens chromosome 21 segment HS21 C085	Home earlies chramosome 21 segment HS210084	III.H.BIS.aks.d-01-0-11.81 NCI CGAP Sub5 Homo sapiens cDNA done IMAGE:2736280 3'	Homo saniens NKG2D gene, exch 10	Truling equipment (CC) and the company of the compa	nom sapisate MAZZZ gate, wor.	COLOR TUTTIBLE I SULLA MACCON HOMEN CON CONTROL IMAGE:3814667 5	6012/20313F1 Nin INGC 20 Total October 1 Angestral handows	Homo saplens Diva, Mino class i region, in directions	Homo saplens mKNA for KIAAU 28 protein, par usa cus	Homo sapiens mRNA for KIAA0768 protein, parties cus	Homo saplens cytoplasmic Seprase fundated isolonii ilinuin, compres exe	
	Top Hit Detabase Source							TO TO	ESI TOMBIA	ESI HUMAN	EST HUMAN	EST HUMAN	L	-	NŢ	<u>.</u>	<u>-</u>	E	Z	Į.	Z	Ł!	Z	Z	Z		NI TOT IS BANK	ESI HOMAN	Z	LN	EST HUMAN	EST HUMAN	LZ.	H	NT	NT.	
	Top Hit Acession No.	11420514INT	TA COLUMN	3		AB020710.1 NT	TIM	1	1				7110834 NT	10634		AF156776.1	11430193 NT	30193	٦		1	AB033104.1	AF084530.1	M30938.1	AL163285.2	AL163285.2	AL163284.2	AW 449746.1	AJ001689.1	AJ001689.1	W26367.1	BE386363.1	AB031007.1	AB018301.1	AB018301.1	AF007822.1	
	Most Similar (Top) Hit BLAST E Velue	1 01.80	3	1.0E-90	1.0E-90/	1.0E-90	10	1.UE-80	8.0E-91	5.0E-91	5.0E-91	5.0E-91	6.0E-91	5.0E-91	4.0E-91	4.0E-91	3.0E-91	3.0E-91	3.0E-91	3.0E-91	3.0E-91	3.0E-91	3.0E-91									8.0E-92	7.0E-92		7.0E-8	7 OF-02	_
-	Expression Signal	2 FEB	37	9.3	0.89	0.99		1.62	6.67	2.28	1.19	1.19	1.09	1.09	1.67	1.67	1.86	1.86	1.4	1.55	3.62	3.62	1.57					6.11	7.33	7.33	89.9	7.89				200	V.9
-	ORF SEQ ID NO:	44000	1820	12819	13765	13766		14278	14057	13428	14355	14356						11638	11830	13301	13410		13706	14428		14786		11271	11265	11266							1
	SEQ ID	200	DCSD	7801	8764	8764			2906	8402	9376	9376	L	1_			L		Ŀ		8388	L	8703	9448	9804	0 9804	9 5130	6225	L		1_		1	١	1		5817
	Probe SEQ ID NO:		1861	2780	3761	3761		4288	4073	3394	4385	4385	4664	4684	3129	3129	1578	1678	1754	3265	3380	3380	3699	4458	4820	4820	49	1226	1222	1222	ā	8	1,	3	8 8	3	586

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo sapiens B-cell CL/Jymphoma 7b (BCL7B) mRNA	Homo sapiens ARP2 (actin-related protein 2, yeast) homolog (ACTR2), mRNA	Homo sapiens ARP2 (actin-related protein 2, yeast) homolog (ACTR2), mRNA	Homo sapiens cysteine-rich repeat-containing protein \$52 precursor, mRNA, complete cds	Homo saplens NRAS-related gane (D1S155E), mRNA	Homo sapiens DNA, MHC class I region, 7.1 ancestral haplotype	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAMI) mRNA	N-CAM=145 kda neural cell adhesion moiscule [human, small cell lung cancer cell line OS2-R, mRNA, 2960 nt	N-CAM=145 kda neural cell adheston molecule [human, small cell lung cancer cell line OS2-R, mRNA, 2960	bu	Homo septients chromosome Z1 segment HSZ1C081	601283012F1 NIH_MGC_44 Hamo sapiens aDNA alone IMAGE:3605018 5	601501242F1 NIH_MGC_70 Hamo sepiens cDNA clans IMAGE:3902939 6	Homo septens activin A receptor, type IIB (ACVR2B) mRNA	Homo sapiens hypothetical protein d.1462023.2 (D.1462023.2), mRNA	Homo sapiens hypothetical protein dJ462O23.2 (DJ462O23.2), mRNA	601118337F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3028304 5		mrg≃mas-related (human, Genomic, 2416 nt)	wk27d07.x1 NCI_CGAP_Briz5 Homo septens cDNA clone IMAGE:2413549.3' similar to TR:Q12844 Q12844 BREAKPOINT CLUSTER REGION PROTEIN;	wk27d07.x1 NCI_CGAP_Bm25 Hano sapiens cDNA clane IMAGE:2413549 3' similar to TR:Q12844	012844 BREAKPOINT CLUSTER REGION PROTEIN;	Homo sapiens syndecan 4 (amphiglycan, ryudocan) (SDC4) mRNA	Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA	Homo sapiens collagen, type XII, alpha 1 (COL12A1), mRNA	Homo sapiens collagen, type XII, alpha 1 (COL12A1), mRNA	Homo sapiens chromosame 21 unknown mRNA	Homo sapiens chromosome 21 unknown mRNA	Homo sepiens stress-Induced-phosphoprotein 1 (Hsp70fHsp90-organizing protein) (STIP1), mRNA	
- -	Top Hit Database Source	F	F	77	NT	7	¥	누	F	_ <b>F</b>	<u> </u>	Z	Z	EST_HUMAN	EST_HUMAN	IN	5	5	EST_HUMAN	EST_HUMAN	47	EST_HUMAN	_	EST_HUMAN	47	F	圬	17	IN	ᅻ	<b>_</b>	NT
	Top Hit Acession No.	4502384 NT	5031570 NT	5031570 NT	F167706.1	4005738 NT	AB031007.1	4507500 NT	4507500 NT			T	1		3.0E-92 BE909714.1	4501898	11422946 NT	11422948 NT	2.0E-92 BE299190.1			1818118.1		1818119.1	4506860 NT	6912457 NT	11418424 NT	8424	2.0E-92 AF231919.1		5803180 NT	10976.1
	Most Similar (Top) Hit BLAST E Value	7.0E-92	7.0E-92	7.0E-92	7.0E-92	7.0E-92		7.0E-92	7.0E-92	7.0E-92 S71824.1	7.05	7.0E-#2.371824.1	/.UE-92 /	6.0E-92	3.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92 A		2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92	2.0E-92 M
	Expression Signal	2.83	8.39	8.39	2.56	5.83	0.93	0.67	0.67	1.24	70.7	1.24	-A-	1.37	2.21	1.42	3.37	3.37	. 2.34	2.34	1.45	1.59		1.59	8.59	15.93	3.89	3.89	1.16	1.16	5.87	1.18
	ORF SEQ ID NO:				12576	12723	12762	13304	13305	14425	80777	44804	2		12764	10090	10251	10252	10787	10788		11976		11977	12084	12666	11876	11677	13543	13544	13617	14147
	Exan SEQ ID NO:	6259							10047	9445	DAAR	8080	200	6549	7850	6106	5241	5241	5763	5763	6672	6884		88	6379	7551	6611	6611	8238	8538	8008	9160
	Probe SEQ ID NO:	1261	2123	2123	2493	2653	2679	3270	3270	4455	4455	4844		1552	2692	92	178	178	740	740	1676	1896		<u>8</u>	1995	2588	2756	2756	3532	3532	3602	4165

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	DKFZp434C0414 r1 434 (syncovym: hiss3) Homo sablens cDNA clove DKF7c434C7444 K	y80e08.r1 Soares placenta Nb24P Homo sepiene cDNA close MAGE-125572 F	y80e08.r1 Soares placenta Nb2HP Homo saciens cDNA clone IMAGE-145574.67	Homo saplens ribosomal protein, large, P1 (RPLP1) mRNA	AU121681 MAMMA1 Hamo septens cDNA clane MAMMA1000738 5	EST188414 HCC cell line (matastasts to liver in mouse) Il Homo septens cDNA 5' end similar to ribosomal protein (.29	601281867F1 NIH MGC 44 Homo saniens citing olivas IMA GENSANSANS E	AU121681 MAMMA1 Homo septems cDNA clone MAMMA1000738 67	Homo saplens chromosome 21 unknown mRNA	Hamo saplens mRNA for KJAA0611 protein, partial cds	wc08c08.xt NCI CGAP Pr28 Homo seplens cDNA clane MAAGE-231487n 31	wc09c08.xt NCI_CGAP_Pr28 Homo eaplens cDNA clone IMAGE:2314870 3'	Human skeletal muscle 1.3 kb mRNA for tropomyosin	260e09.s1 Sogres_lestis_NHT Homo saplens cDNA clone IMAGE:785688 3' similar to SW:CLPA_RAT P37397 CALPONIN, ACIDIC ISOFORM:	Homo sapiens interferon gamma receptor 1 (IFNGR1) mRNA	Homo saplens interferon gamma receptor 1 (IFNGR1) mRNA	Homo sapiens pescadillo (zebrafish) homdog 1, containing BRCT domain (PES1), mRNA	Homo sapiens pescadillo (zebrafish) homolog 1, containing BRCT domain (PES1), mRNA	Homo saplens hypothetical protein FLJ20731 (FLJ20731), mRNA	Homo saplens dystrophin (DMD) gene, deletion breakpoints 1-3 in intron 5	Homo sapiens DNA polymerase zeta catalytic subunit (REV3) mRNA, complete ods	Homo septens TNF-Inducible protein CG12-1 (CG12-1), mRNA	Homo sapiens fumor antigen SLP-8p (HCC8), mRNA	Homo sapiens Interfeukin 18 receptor 1 (IL18R1) mRNA	Homo saplens tumor antigen SLP-8p (HCCs), mRNA	602246554F1 NIH_MGC_82 Homo sapiens cDNA clane IMAGE:4332038 67	602246554F1 NIH_MGC_62 Homo sapiens cDNA clone IMAGE:4332036 5	Chlorocebus aethiops mRNA for ribosomal protein S4X, complets cds	Chlorocebus aethicps mRNA for ribosomal protein S4X, complete cds	Homo saplens chromosome 21 segment HS21C085	Homo seplens chromosome 21 segment HS21C085
	Top Hit Database Source	EST HUMAN	EST HUMAN	EST HUMAN		LHUMAN	EST HUMAN	EST HUMAN	Г	Г	1	EST HUMAN	Г		EST HUMAN	_	1	Þ	_	1	Ţ				Ţ	T		T_HUMAN	1			
,	Top Hit Acession No.	L040437.1	78078.1	Γ	450668 NT	9.0E-83 AU121681.1 E	9.0E-83 AA316723.1	Γ	Ī	Γ	Γ				AA459833.1 E	57879	4557879 NT	7657454 NT	7657454 NT	8923658 NT	047677.1	F157476.1 NT	7656972 NT	7705396 NT	4504654 NT	7705396 NT	3.0E-93 BF690630.1			8015610.1		2.0E-93 AL163285.2 INT
	Most Similar (Top) Hit BLAST E Value	2.0E-92 A	1.0E-92 R78078.1	1.0E-92 R78078.1	1.0E-92	9.0E-83	9.0E-93.A	9.0E-93 B	9.0E-93 A	7.0E-93 A	6.0E-93	6.0E-93 AI674184.1	5.0E-93 A	5.0E-93 X04201.1	4.0E-93 A	4.0E-93	4.0E-93	4.0E-93	4.0E-93	4.0E-93	4.0E-93 AF	≤।	4.0E-93	4.0E-93	4.0E-93	4.0E-93	3.0E-93 B	3.0E-93 B	2.0E-93 A	2.0E-93 A	2.0E-93 A	2.0E-93
	Expression Signal	2.79	2.03	2.03	40.83	2.63	27.81	1.75	1.1	8.34	2.07	8.53	8.53	4.58	5.69	1.62	1.62	4.03	4.03	1.25	3.59	0.93	1.01	0.79	5.14	0.83	19.66	19.66	31.68	31.68	9.39	6.30
	ORF SEQ ID NO:		11895	11896	12108	12070		13546	14184	10314	11409	11429	11430	13200		10496	10497	10812	10813	11201	12017	12282	12618	13512	13927	13512	13579	13580	10265	10266	10386	10386
	Exan SEQ ID NO:			6803	7003	6964	6876	8540		5304		6380	6380	8178	5163	5479	5479	6784	5784				$_{ m l}$	8495	8334	8495	8574	8574	5254	5254	5376	6376
	Probe SEQ ID NO:	4835	1813	1813	2020	1979	1991	3534	4209	244	1362	1383	1383	3162	86	442	442	783	763	1164	1932	2183	2533	3487	3935	4863	3567	3567	190	<u>19</u>	321	322

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Too Hit Descriptor		601117686F1 NIH MGC 16 Homo seniens CDNA close (MACE 198822) R	801116810F1 NIH MGC 16 Hamo septens cDNA clans IMAGE 3357923 6	Homo saplans CTR1 pseudopana	Homo saplens CTR1 pseudocene	Homo saplens hypothetical protein (DJ328E19.C1.1), mRNA	oyekbo8.x1 NCI_CGAP_CL11 Homo sepiens cDNA clane IMAGE:1672503 3' similier to TR:062384 Q62384 I ZNC FINGER PROTFIN	Homo sablens DNA for amyloid precursor protein committee of a	Homo saplens hypothetical protein FL 120201 (FL 120201) mRNA	Homo sapiens hypothetical protein FL 20291 (FL 20291) mRNA	Homo sepiens mRNA for KIAA1563 protein, partial cds	Homo sapiens cysteine-rich repeat-containing protein S52 precursor, mRNA, complete cds	Homo sabiens long chain bolyunsaturated fatty and elongation entering (HFI 0.1) and About	Homo saplens MHC class 1 recton	Novel human gene mapping to champagne 1	601177686F1 NIH MGC 17 Hamo saniers cDNA clone IMAGE 3533264 F	801177686F1 NIH MGC 17 Hamo sablens cDNA clane IMAGE:3532965 F	Homo sapiens DNA for amyloid precursor protein, complete cds	Homo saptens tong chain polyunsaturated fatty acid elongation enzyme (HFI O1) mBNA complete ade	Hamo sapiens chromosome 21 segment HS21C084	Homo sapiens transcription enhancer factor-5 mRNA, complete cds	Homo sapiens ribosomal protein L27 mRNA, complete cds	Homo sapiens protein phosphatase 1, regulatory subunit 10 (PPP1R10) mRNA	m89f12.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA done IMAGE:2701679 3'	xn89f12.x1 Sogres_NFL_T_GBC_S1 Hamo septens cDNA clane IMAGE:2701679.3'	tw11f10.x1 NCI_CGAP_Bm52 Hamo saplens cDNA clone IMAGE:2258403 3' similar to TR:Q15265 Q15285	PROTEIN TYROSINE PHOSPHATASE; Homo santens ASH21 nens commiste of selections to the selections of the commister of selections and the selections of the commister of selections and the selections are commistered to the selections and the selections are commistered to the selections and the selections are commistered to the selections are commissed t	Home sablens combinent component 5 (C5) mRNA	Homo saplans cysteine-rich rapeat-containing protein SS2 precurear, mRNA complete ode	Homo saplens cysteine-rich repeat-containing protein S52 precursor, mRNA, complete cds	Homo saplens E1A binding protein p300 (EP300) mRNA	
	Top Hit Database	Source	EST HUMAN	EST HUMAN				EST HUMAN									EST HUMAN	EST HUMAN							EST_HUMAN	EST_HUMAN		EST HUMAN					
L			EST	EST	E	ż	Ż	EST	ż	Ż	눌	Ę	'n	Ë	ż	ż	EST	EST	Ę	ż	눌	Ę	Þ	LΝ	EST	EST		E E	Ę	Ę	Ę	μ	
	Top Hit Acession	o Z	BE252982.1	2.0E-03 BE263201.1	1.0E-93 AF238997.1	1.0E-93 AF238997.1	7857018 NT	1.0E-93 Al146755.1	D87675.1	8923270 NT	l	1.0E-93 AB046783.1	1.0E-93 AF167706.1	1.0E-93 AF231981.1	1.0E-83 AF055066.1	1.0E-93 AL137200.1	1.0E-93 BE297369.1	1.0E-93 BE297369.1	1:0E-83 D87675.1	1.0E- <del>0</del> 3 AF231981.1	1.0E-03 AL163284.2	6.0E-84 AF142482.1	.05094.1	4506008 NT	4W197851.1	4.0E-94 AW197851.1		4.0E-94 AI591312.1 3.0F-94 AR022785 1	4502508 NT	18	3.0E-94 AF187708.1	4557556 NT	
	Most Similar (Top) Hit	Value	2.0E-83	2.0E-03	1.0E-93	1.0E-93	1.0E-93	1.0E-83	1.0E-03 D8767	1.0E-93	1.0E-93	1.0E-93	1.0E-93	1.0E-83	1.0E-93	1.0E-93	1.0E-93	1.0E-93	1:0E-93	1.0E-93	1.0E-93	6.0E-94	4.0E-84 L050	4.0E-94	4.0E-94 AW1	4.0E-94	-	3 0F-94 AR02	3.0E-04	3.0E-94 AF16	3.0E-94	3.0E-94	
	Expression		1.74	1.01	2.66	2.66	17.48	3.67	7.39	8.15	8.15	1.13	3.03	6.14	1291	1.31	1.38	1.39	3.34	1.76	2.36	3.44	38.53	1.13	1.02	1.02		3.55	1.52	3,56	3.56	6.42	
	ORF SEQ	<u></u>	12507	14920	10189	10190	10554	10625	10921	. 11256	11257	11370	11372	12370	12487		11316	11316	12903		14283	13864		12664	13598	13599		10635	10751	11772	11773	11807	·.
	Exan SEQ ID	ğ	7387		5179	6179	5549			6216		6323	6325	7252	7365	7408	6275	6275	7883	8160	9297	8858	6797	7549	8594	8594		9567	6735	9699	9699	6730	
	Probe SEQ ID	Ö	2418	4966	102	102	514	595	198	1217	1217	1325	1327	2276	2394	2435	2749	2749	2863	3144	4305	3856	1806	2586	3587	3587	į	49/9	711	1701	1701	1735	

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Single Exon Probes Expressed in HBL100 Cells	Top Hit Descriptor	2w63g08.r1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:774782.6	Homo saplens ubiquitin specific protease 13 (Isopentidase T.3) (USP 3) mRNA	601175762F1 NIH_MGC_17 Hamo suplens cDNA clane IMAGE:3531038 5	601111696F1 NIH_MGC_16 Homo septens cDNA clone [MAGE:3352559 57	601111696F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3352559 67	Homo sapiens hypothetical protein (FLJ20748), mRNA	Homo saplens TNF-alpha stimulated ABC protein (ABC50) mRNA, complete cds	Homo saplens KIAA0255 gene product (KIAA0255), mRNA	Homo sapiens KiAA0255 gene product (KIAA0255), mRNA	we08e04.x1 NCI_CGAP_Lu24 Homo sepiens cDNA clone IMAGE:2340608 3' similar to gb:K00568 TUBULIN ALPHA-1 CHAIN (HUMAN);	we09e04x1 NCL_CGAP_Lu24 Homo saplens cDNA clone IMAGE:2340606 3' similar to gb:K00558 TUBULIN ALPHA-1 CHAIN (HUMAN):	Homo seplens DNA for amyloid precursor protein, complete cds	Homo sapiens DNA for amyloid precursor protein, complete cds	Homo sapiens Ly-6-like protein (CD59) mRNA, complete cds	Homo saplens chromosome 21 segment HS21C046	Homo sapiens KIAA0255 gane product (KIAA0255), mRNA	Homo sapiens KIAA0255 gene product (KIAA0255), mRNA	Homo sepiens tissue inhibitor of metalloproteinase 3 (Sorsby fundus dystrophy, pseudoinflammatory) (TIMP3) mRNA	601312161F1 NIH_MGC_44 Homo saplens cDNA clone IMAGE:3658862 6'	Homo saplens G protein-coupled receptor 19 (GPR19) mRNA	Homo saplens G protein-coupled receptor 19 (GPR19) mRNA	Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1)	Homo saplens glycine cleavage system protein H (aminomethyl carrier) (GCSH) mRNA	Homo sapiens Usurpin-gamma mRNA, complete cds	Homo sapiens unconventional myosin-15 (LOC51168), mRNA	Homo sapiens unconventional myosin-15 (LOC51168), mRNA	Homo sapiens mRNA for KIAA1386 protein, partial cds	qm01c02.x1 Soares_NhHMPu_S1 Homo sapiens cDNA clone IMAQE:1880546 3' similar to WP:123G7.4 CE03705;	Homo sapiens hypothetical protein (HS322B1A), mRNA	
sedory no	Top Hit Database Source	EST_HUMAN	1	EST HUMAN	EST_HUMAN	EST_HUMAN	E	ᄫ	E	H	EST_HUMAN	EST HUMAN	Į.	N	IN	NT	E	<u> </u>	<u> </u>	EST_HUMAN	Τ	T	5		IN.	E	1	NT	EST_HUMAN	E	_
Ca eignic	Top Hit Acession No.	AA464805.1	4507848 NT	BE295714.1 E	BE253433.1 E	BE253433.1 E	26692	AF027302.1	7662027 NT	7662027 NT	Al700998.1	AI700998.1 E	D87675.1	D87675.1	M95708.1	AL163246.2 N	7862027 NT	7662027 NT	4507512 NT	BE393873.1 E	5453665 NT	5453665 NT	AF240788 1	38423	AF015452.1	TN 0065077	7705900 NT	AB037807.1	A1290264.1	7657185 NT	
	Most Similar (Top) Hit BLAST E Value	3.0E-84	3.0E-04	1.0E-94	1.0E-94	1.0E-94	1.0E-94	9.0E-95	9.0E-95	9.0E-95	8.0E-95	8.0E-95	7.0E-95	7.0E-95	7.0E-95	7.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	20.40	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	2.0E-95	
	Expression Signal	0.67	0.7	2.88	2.59	2.59	1.7	5.95	1.09	1.09	3.37	3.37	10.63	10.63	5.66	1.4	3.13	3.13	3.11	1.74	1.55	1.65	2 79	1.84	1.85	2.78	2.78	1.17	1	1.61	
	ORF SEQ ID NO:	14048			13042		14211	11503	13112	13113	14380	14381	10344	10345	14219		11662	11663	11982	11986	12449	12450	12488	12528	13115	13508	13509	13549	13674	14213	
	Exan SEQ ID NO:	6906	10022		8032		9227	6444	8608	8098	8397	9397	5332	6332	9235	9281		6601	6888	6891	7333	7333	7367	7412	8100	8492	8492	8543	8669	9230	
	Probe SEQ ID NO:	4085	5051	148	3015	3015	4233	1447	3082	3082	4406	4406	273	273	4241	4289	1605	1605	1901	1904	2359	2359	9682	2442	3084	3484	3484	3537	3664	4236	

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo senjens KIAA0187 gene product (KIAA0187), mRNA	TALLES TO THE TALL NICHES ON Home samens CONA clone IMAGE: 786167 6	2X1100/.11 SORIOS COM 1000 1 TO 1 TO 1 TO 1 TO 1 TO 1 TO 1 TO	EXTIGO/ IT SOME BUILD IN THE SOUTH AND INTEREST SOUTH FOR	601497608F1 NIH MIGG 70 Hamp septients Chira clare impost and a septients chira clare and a septients chira clare and a septients chira clare and a septients chira clare and a septients chira clare and a septients chira clare and a septients chira clare and a septients chira clare and a septients chira clare and a septients chira chir	601497608F1 NIH_MGC_/0 Homb september curva cum invacingos	Home saplens chromosome 21 unknown mrtvA	MRO-HT0559-250200-002-407 H10599 Homo septems cures	Homo sapiens chromosome 21 segment no 210001	Human glyceraldehyde-3-phosphate denydrogenase pseudogene s end	Homo saplens mKNA 10' KIAA11/2 protein, per use vus	Homo seplens mRNA for KIAA11/2 protein, partial cus	Home sapiens mRNA for KIAA1172 protein, partial cos	Homo seplens phosphodiesterase 6A, cGMP-specific, rod, alpha (PDE5A), mytwa	H. sapiens DNA for monoamine oxidase type A (7) (partial)	yr87h12.r1 Sogres fetal liver spleen 1NFLS Home saplens CDNA clone IMAGE: 212327 o	Home sepiens chondraltin sulfate proteoglycan 4 (melanoma-associated) (CSF C4), minum	Homo sapiens chromosome 21 segment HS21C048	Homo eaplens CGI-201 protein (LOC51340), mRNA	RC3-HT0230-040500-110-d02 HT0230 Homo sapiens cDNA	Human endoremus retrovirus type K (HERV-K), gag, pol and env genes	FECTABA12A MAGE resentances. MAGC Homo saplens cDNA	EST367124 MAGE resemences, MAGG Homo saplens cDNA	E-its able simplest more in the save their (sMVHC) mRNA. complete cds	For Subsider 4 70300-203-408 RN0106 Home septens CDNA	Christian Control of the Publish of Christian Publish of Christian (BIG2), mRNA	Florito Septens Distriction of Transfer and	HOMO Sapiens minny lot Nich in Fromin, principal	Homo sapiens amyloid bets (A4) precursor protein (protesse nextr-II, Abrhelmer disease) (APP), mRNA	Homo saplens amykold beta (A4) precursor protein (probase nextn-ii, Aizheimer disease) (APP), mRNA	Homo saplens N-myc (and STAT) Interactor (NMI), mRNA	Hirman beta-orime-edaptin (BAM22) gene, exon 7	Home seciens percentin (PCNT) mRNA	Homo senians eukanotic translation elongetion factor 1 alpha 1 (EEF1A1) mRNA	
	Top Hit Database Source			EST HUMAN	EST_HUMAN	EST HUMAN	EST HUMAN	NT -	EST_HUMAN	N <sub>T</sub>	N	L	1	NT	1	NT	EST_HUMAN	<u> </u>	LZ	L	FOT LIMAN	L L	N HOL	ESI FORMAN	ESI HOMAIN	IN.	EST HUMAN		L	, 5	<u>F</u>	5		E F	E P	
218110	Top Hit Acession No.	7884070 MT		7						163201.2		5.0E-96 AB032998.1 NT	5.0E-96 AB032998.1 NT		11416787 NT			4503098 NT	2 OF 06 AI 183248 2	8005		E148074.1	18890.1	T	-		BE004436.1	3972	3.0E-97 AB032998.1	4502166 NT	. 4502186 NT	TN 619844	210001	U36255.1		4503470141
-	Most Similar (Top) Hit BLAST E Value	9000	Z.UC-83	2.0E-85 AA447831.1	2.0E-95 AA447831.1	8.0E-96 BE907607.1	8.0E-96 BE907607.1	7.0E-96 AF231920.1	6.0E-96 B	8.0E-96 AL	6.0E-96 M26873.1	5.0E-96 A	5.0E-96 A	5.0E-96	5.0E-98	5.0E-96 X60812.1	3.0E-96 H68656.1	2 0F-96	20506	2000	Z.UE-90	2.0E-96 t	1.0E-96 Y	1.0E-96 /	1.0E-96/	1.0E-96	4.0E-97	4.0E-97	3.0E-97	3.0E-97	١.					1.08-8/1
-	Expression Signal		200	. 0.92	. 0.92	3.42	3.42	1.16	2.4	0.86	37.31	3.23	3.33	3.33	2.15	1.28	10.55	4 94	10.	7. 7.						1.95		1.34	2.28	9.6						35.24
	ORF SEQ ID NO:		14827	14869	14870	10483	10494	13822	12299	13278	13437			10890										11814	11815	12305	10976	11944	10312	10923						14612
	Exan SEQ ID NO:		9852	9895	9895	7720	7720	8816	7176	8257			L	L			_	l	_					6738	6738	17705	5 5942	7 6856		5882				1 7706	6 8202	5 9620
	Probe SEQ ID NO:		4873	4917	4917	439	439	3813	2198	3244	3402	318	832	1	2545	36	4740	200	412	738	1756	4613	663	1743	1743	2204	925	1867	241	983	8	863	1415	2371	3186	4635

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	PLAA BT0724-010400-008-s12 BT0724 Homo sapiens cDNA	The second section of the second second (CEOR1) INRINA	Homo seguests can by entrangent (18PR1 ref.) homolog 1 (FSHPRH1), mRNA	JOHO SEPTEMBER TOT METERS FROM LONG THE PROPERTY AND THE	Homo sapiens partial MICB gene for MIHC class I chain-related protein B, exons 2-3 and johned CDS	Homo sapiens IL2-inducible T-cell kinase (ITK), mRNA	Homo sapiens II.2-inducible T-cell kinase (ITK), mittina	Homo septens PMSZL16 mKNA, perus cas	Homo sapiens PMSZ.16 mKNA, partial cos	Human mitochondrial creature kinase (Orwit) gains, compress con 2000 clone 18	AJ403124 S.4 (downregulated in lat pit concurrent) forms of the control of the co	HOMO Sapletts MINNA IN NAVA OF Presh CDNA I Trany Homo saplens CDNA Clone 7B18H01	ASTACHOL CHAMING INC. 10 Home entires CDNA clane IMAGE:3502245 6	601149460F1 NIT MCC, 19 Holling saylans CDNA clone IMAGE:3628134 5	001112030FT NILL MODEL TO THE PROPERTY OF THE	Homo saprens durinosante z l'ocurrente de l'acceptante de l'ac	Home enjage indesting channel subunit (HERG-3) mRNA, complete ods	Umma contants (ath-eoid-Coenzyme A ligase, long-chain 4 (FACL4) mRNA	Home saniens attractin precursor (ATRN) gene, exen 16	Homo saniens attractin precursor (ATRN) gene, exon 16	1. DANA Y NOT CRAP UM Homo sapiens CDNA clone IMAGE: 2281743 3' similar to SW:RL2B_HUMAN	P28316 60S RIBOSOMAL PROTEIN L23A ;	PM0-BN0065-100300-001-c06 BN00000 Floring september CDNA clone IMAGE-243685 5 similar to	y/23/05.r1 Soares teta tive spicen tiving appearation of the second section of the second second protein L29 - human;	Harbor explains cystaine-rich repeat-containing protein S52 precursor, (LOC61232), mRNA	Truing explains cretains rich reneal-containing protein S52 precursor, (LOC51232), mRNA	Litera angles Cyan antinen (CD34) mRNA	House septens Coort aring at 1	Linguistical International Para locus, TCRBV7S3A2 to TCRBV12S2 region	Home seriens ubjaulitie-contrasting BIR-domain enzyme APOLLON mRNA, complete cds	Home september thin utility conjugating BIR-domain enzyme APOLL ON mRNA, complete cds	Thirty captures are a grant of the control of the c
	Top Hit Datebase Source	Т	NO POR			NT.	Ш	T	NT	۲.	7	EST HUMAN	LV.	EST HUMAN	EST HUMAN	EST HUMAN	Ł !		ź	Z	12	Z	EST_HUMAN	EST HUMAN		EST HUMAN	ž	Z.	Ž.	z!	ŁN.	Į.	N.
,	p Hit Acession No.		JEOS/3.1	8393092 NT	11419594 NT	251168.1	5031810	5031810 NT							1		2.0E-98 AL163202.2	88	2.0E-98 AF032897.1	8331	T	2.0E-98 AF218902.1	862007.1	1.0E-98 AW998611.1	- 10	188 188 188	11430555 N.I	11430555INI	4502660 NT		<b>∢</b> I		AF265555.1
-	Most Similar (Top) Hit BLAST E		9.0E-98 BE	9.0E-98	9.0E-98	8.0E-98 AJ	8.0E-98	8.0E-98	8.0E-98 AB017007.1	8.0E-98 AB017007.1	8.0E-98 J04469.1	3.0E-98 AJ	3.0E-98 A	3.0E-98 A	2.0E-98 B	2.0E-98 B	2.0E-98 A	2.0E-98	2.0E-98 A	2.0E-98	2.0E-98 /	2.0E-98 /	1.0E-98 AI					ı					6.0E-99
	Expression Signal	-	6.71	1.1	. 1.03	4.32	108	1.06	3.31	3.31	6.04	1.08	1.4	2.13	2.43	2.25	2.4	0.94				1.61	91.85			68.95	1.6	1.5		5 0.92	1.25	1.82	1.82
	ORF SEQ ID NO:		10948	11297	14886		44584					12207	12619		10770	12115	12275	13973	14154		14846	14647	10450	١		3 11838	12159	12160		12005		14584	14565
	Exan SEQ ID NO:		8063	8255	9992	8404	9104 8628	6526		L	L				L	7009	L	8987	9167	9207	9996	9998	6430		上	1 6753	7051	L	L		L	L	Ш
		_	98	1257	5021		\$ 8	200	18	1688	3706	2113	2534	2678	728	2026	2176	3989	4172	4214	4680	4680	1 8	3 5	šÌ	1761	2069	2069	4605	1924	1122	4586	4586

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Describior		XXXXXXXI NCI_CGAP_HN9 Homo septens cDNA done INAGE:2739874 3' similar to gb:N31212 MYOSIN LIGHT CHAIN ALKALI NON-ARISCI E ISOFORM (HI IMAN):	Human Ku (p70/p80) subunit mRNA, complete cds	Homo seplens short chain L-3-hydroxyecyl-CoA dehydrogenase precursor (HADHSC) gene, nuclear gene encoding mitochandrial protein, complete cds.	Homo septents Intersectin long (soform (ITSN) mRNA, complete ods	Homo sapiens GA-binding protein transcription factor, alpha subunit (60kD) (GABPA), mRNA	Human Ku (p70/p80) subunit mRNA, complete cds	Homo septens fruncated Nemarm-Pick C3 protein (NPC3) mRNA, complete ods	Hamo sapiens transated Niemann-Pick C3 protein (NPC3) mRNA, complete cds	Homo saplens PK508-binding protein 6 (38/Q) (FKBP8) mRNA, and translated products	Homo saplens FK508-binding protein 6 (36kD) (FKBP6) mRNA, and translated products	Human Interferon-alpha receptor (HulFN-alpha-Rec) mRNA, complete cds	Homo sapiens fatty acid amide hydrolase (FAAH) gane, exon 14	Homo sapiens fatty acid amide hydrolase (FAAH) gene, exon 14	Homo saplens chromosome 21 segment HS21C047	Homo sapiens chromosome 21 segment HS21C047	Homo saplens Testis-specific XK-related protein on Y (XKRY), inRNA	Homo sapiens Testis-specific XK-related protein on Y (XKRY), mRNA	x/78b11.x1 NCI_CGAP_Bm53 Homo septens cDNA clone IMAGE:2824605 3*	Homo saplens chromosome 21 segment HS21C006	Homo sapiens chromosome 21 segment HS21C049	EST02975 Fetal brain, Stratagene (cat#838208) Homo sapiens cDNA clone HFBCR32	apiens X-linked anhidrotito ectodermal dysplasta protain gene (EDA), exon 2 and flanking repeat		G.gorilia DNA for ZNF80 gene homolog	RC3-HT0825-040500-022-b09 HT0625 Homo septens cDNA	Homo sapiens DKFZP688M0122 protein (DKFZP686M0122), mRNA	Homo sapiens DKFZP586M0122 protein (DKFZP586M0122), mRNA	UI-H-BI1-effk-c-07-0-UI,s1 NCI_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2722164 3'	q/62/09.x1 Soeres_testis_NHT Homo saplens cDNA done IMAGE:1764633 3' stmiler to SW:CYT_COTJA	Homo cantana mRNA for KIA 64468 mental ade	TOURS SEPTION IN TAILY IN THE TOP HE CASE.	nome sapiens nidaluse/ protein (nidaluse/), micha
Top Hit Detabase	Source	EST HUMAN	F	<u> </u>	¥	7	N	Ę	N	NT	=	NT.	LN LN	NT	N	N	T)	J.,	EST_HUMAN	TN	NT	EST_HUMAN		NT	L	EST_HUMAN	11	<u> </u>	EST_HUMAN	COT LIMIAN	TO TOWN		
Top Hit Acession		AW274792.1		AF095703.1	AF114487.1	11526150 NT	M30938.1	AF192523.1	AF192523.1	4503730 N	4503730 NT	J03171.1	AF098018.1		AL163247.2 IN		11418230 NT	11418230 NT	AW275237.1	AL163206.2 N	AL163249.2	T05087.1		-		BE180609.1 E	7681685 NT	7861685 NT	AW207555.1	A12000874	T	92.00	11418976 N
Most Similar (Top) Hit	Value	2.0E-99	2.0E-99	2.0E-89	1.0E-99	1.0E-89	1.0E-99	1.05-99	1.0E-99	1.0E-99	1.0E-99	1.0E-99	1.0E-99	1.0E-99	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100		1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	1.0E-100	4 05 400	201-101	1.001-301	1.UE-10VI
Expression		21.29	2.08	. 2.65	1.21	1.23	8.91	3.27	3.27	1.12	1.12	1.27	2.45	2.45	1.69	1.64	1.24	1.24	2.54	0.69	1.05	1.8		1.98	19.01	1.38	2.48	2.48	1.64	47 7	2 4	F	ZC.I.
ORF SEQ			13223	14387	10379	10440			11579		11965	13039	14233		10067	10067	10151	10152	10174	10243	10381	10400				10547	11042	11043		44579	1973	11811	12/0/1
Exan SEQ ID	ö	6220	8200	9402		6425			6522	6875	6875	8028	8248	9248	5083	5083	5146	5146	5162	5233	5370	5393		8472	6523	5541	6013	6013	6512	9139	514	3 3	100
Probe SEQ ID	ö	1220	3184	4412	313	378	1396	1525	1525	1886	1886	3011	4255	4255	•	2	67	67	85	168	315	341		\$	\$	909	1003	1003	1514	4640	1827	700	ğ

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Homo sapiens RGH2 gene, retrovirus like element	Homo sepiens myotubularin-related protein 1a mRNA, partial cds	Homo saplens follicle stimulating hormone receptor (FSHR) mRNA	Homo sapiens small optic lobes (Drosophila) homolog (SOLH) mRNA	Homo sapiens small optic lobes (Drosophila) homolog (SOLH) mRNA	Homo sapiens SEC14 (S. cerevísiae)-like 2 (SEC14L2), mRNA	Homo sapiens SEC14 (S. cerevisiae)-III/e 2 (SEC14L2), mRNA	Homo sapiens mRNA for KIAA0446 protein, partial cds	Homo sapiens ventral anterior homeobox 2 (VAX2), mRNA	Homo sapiens ventral enterfor homeobox 2 (VAX2), mRNA	Homo sapiens pescadilio (zebrafish) homolog 1, containing BRCT domain (PES1), mRNA	Homo sapiens phosphoribos//djycinamide formytransferase, phosphoribos//djycinamide synthetase,	prospinancosysteminatable symmetase (GART) mRNA	OUZION+/4FI NIN MCC_SO HOMO SEPENS CONA CIONE IMAGE:428/281 5	qg98e09.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:18433363'	Homo sapiens butyrophilin, subfamily 2, member A1 (BTN2A1), mRNA	Homo saplens butyrophilin, subfamily 2, member A1 (BTN2A1), mRNA	Homo sapiens KIAA0569 gene product (KIAA0569), mRNA	Hamo sapiens KiAA0569 gene product (KIAA0569), mRNA	Homo sapiens carboxypeptidase A1 (pancreatic) (CPA1) mRNA	RC3-ST0281-160600-016-h09 ST0281 Homo sepiens cDNA	Homo sapiens A kinasa (PRKA) anchor protein 6 (AKAP6), mRNA	H.sapiens EWS gene, exon 5	Homo saplens RiBilR gene (partial), exon 12	Homo sapiens RIBIIR gene (partial), expn 12	Homo sapiens genomic downstream Rhesus box .	Homo sapiens gamma-glutamyltransferase 1 (GGT1) mRNA	601458531F1 NIH_MGC_68 Homo saplens cDNA clone IMAGE:3882086 5'	EST377629 MAGE resequences, MAGI Homo saplens cDNA	Homo sapiens RIBIIR gene (partial), expn 12	Homo sapiens RIBIIR gene (partial), exon 12	Homo sapiens ASH2L gene, complete cds, similar to Drosophila ash2 gene	Homo sapiens butyrophilin, subfamily 2, member A1 (BTN2A1), mRNA	Homo sapiens butyrophilin, subfamily 2, member A1 (BTN2A1), mRNA	
Top Hit Database Source	¥	¥	5	7	5	5	5	N	=	5	17	!	- 16	ш	EST_HUMAN	=	4	4	11	לד	EST_HUMAN	5	NT	NT	NT	NT	47	EST_HUMAN	EST_HUMAN	N P	LN LN	N	K	47	
Top Hit Acessian Na.	78.1	77364.1	4503782	5032104 NT	5032104 NT	7110714 NT	7110714 NT	07915.2	7110734 NT	7110734 NT	7657454 NT		41851	1.0E-101 DF001210.1		5921460 NT	5921460 NT	7662183 NT	7662183 NT	4502996 NT	43070.1	5729862 NT		1.0E-101 AJ237744.1	1.0E-101 AJ237744.1	1.0E-101 AJ262312.1	4885270 NT	1.0E-101 BF035327.1 E	985558.1	1.0E-101 AJ237744.1		22785.1	5921460 N	5921460 NT	
Most Similar (Top) Hit BLAST E Value	1.0E-100	1.0E-100 AFO	1.0E-100	1.0E-100	1.0E-100	1.0E-101	1.0E-101	1.0E-101 AB0	1.0E-101	1.0E-101	1.0E-101	707	1.0E-101	101-301	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101 BE8	1.0E-101	1.0E-101 X72993.1	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101	1.0E-101 AW	1.0E-101	1.0E-101	1.0E-101 ABC	1.0E-101	1.0E-101	
Expression Signal	4.16	1.67	228	3.82	3.82	1.88	1.88	2.59	6.32	6.32	4.64	900	37.36	00:40	1.9	207	2.07	1	1	1.32	261	76.0	6.73	5.15	5.15	12.51	2.4	2.76	2.01	3.42	3.42	5.05	1.61	1.61	
ORF SEQ ID NO:		14074	14093	14872	14873	10164	10165	10710	10735	10736	10811	40002	14020	201	11074	11606	11607	11779	. 11780	11987	12090	12381	12625	12741	12742		13167		13332	12741	12742	13792	14823	14824	
SEQ ID	7971	9083	9108	8888	9888	5154	6154	5702	5718	5718	5783	6083	3002 FORF	3 5	6045	6546	6546	6702	6702	6892	9869	7767	7506	7628	7628	7903	8146	8183	8307	7628	7628	8788	9847	9847	
Probe SEQ ID NO:	292	4089	4114	4920	4920	9/	92	229	694	694	762	0.42	070		1035	- 25 25 25 26 26 26 26 26 26 26 26 26 26 26 26 26	15 <u>48</u>	1707	1707	1905	2003	2288	2541	2670	2870	2884	3130	3167	3296	3316	3316	3785	4868	4868	

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Top Hit Descriptor	Homo sapiens phosphatidylinosital 4-kinase 230 (p4K230) mRNA, complete cds	Homo saplens chromosome 21 segment HS21C103	601108292F1 NIH_MGC_16 Homo sapiens cDNA clane IMAGE:3344328 5	Homo sapiens down-regulated in adenoma (DRA) mRNA	Human endogenous retroviral DNA (4-1), complete retroviral segment	Homo saplens solute carrier family 2 (facilitated glucose transporter), member 9 (SLC2A9), mRNA	Homo sapiens solute carrier family 2 (facilitated glucose transporter), member 9 (SLC2A9), mRNA	601289982F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3628901 5'	am60c10.x1 Johnston frontal cortex Homo sapiens cDNA clone IMAGE:1539854 3' struitar to SW:GG95_HUMAN Q08379 GOLGIN-95. ;	am60c10.x1 Johnston frontal cortex Homo sapiens cDNA clone IMAGE:1539954 3' almilar to SW:GG85_HUMAN Q08379 GOLGIN-95; ;	Homo sapiens peradsome blogenesis factor 1 (PEX1), mRNA	Homo sapiens KIAA0187 gene product (KIAA0187), mRNA	AU141005 PLACE4 Homo sapiens cDNA clone PLACE4000650 5	AU141005 PLACE4 Hamo sepiens aDNA alone PLACE4000650 6	Homo saplens chromosome 21 segment HS21C007	601107843F1 NIH_MGC_16 Homo sapiens cDNA clane IMAGE:3343882 6	y/32c04.r1 Soares placenta Nb2HP Homo sapiens cDNA clone IMAGE:140934 o	601500405F1 NIH_MGC_70 Homo sapiens cDINA clone IMAGE:3902305 b	601500405F1 NIH_MGC_70 Homo sapiens cDNA ctone IMAGE:3902305 5	Homo saplens mRNA for KIAA0235 protein, partial cds	Homo sapiens nucleolar protein (KKE/D repeat) (NOP-56) mKNA	Homo sapiens mRNA for pregnancy-associated plasma protein-E (FAFTE gand)	601485388F1 NIH MGC 69 Homo sapiens CUNA clone IMAGE:3887670 3	Homo sapiens phosphalayanositol 4-knase 230 (planz3u) mrana, compress cas	Homo sapiens and GLX-AXXOCIATED PROTEIN (WAAP), MINNA	Homo sapiens bone morphogenetic protein 8 (osteogenic protein 2) (BMPs) mixiva	Homo saplens bone morphogenetic protein 8 (ostaogenic protein 2) (brint 8) micros	AU134991 PLACE1 Homo sapiens cDNA clone PLACE1000965 5	Homo sapiens promyelocytic leukemia zinc finger protein (PLZF) gene, complete cas	yw91d08.s1 Soares_placenta_8tc9weeks_2NbHP8to8W Homo sepiens cDNA clone IMAGE:268699 S*	601573113F1 NIH_MGC_9 Homo sapiens cDNA clore IMAGE:3834315 5
Top Hit Database Source			THUMAN					EST_HUMAN	EST_HUMAN	EST_HUMAN			EST_HUMAN	EST_HUMAN	ı	EST HUMAN	T HUMAN	EST HUMAN	EST HUMAN	ٺ	٠	1	EST HUMAN		٠		<b>-</b>	EST_HUMAN	1	EST_HUMAN	EST HUMAN
Top Hit Acession No.	F012872.1 NT	L163303.2 NT	E252470.1 EST	4557534 NT	110976.1 NT	11437146 NT	11437146 NT	8447.1	669.1	1869.1	11419442 NT	TN 6761937	41005.1	11005.1	3207.2	51310.1	188.1	38158.1	-	387078.2 NT:	5453783	٦	77541.1	<u>8</u>	7657592 NT	4502428 NT	4502428	34991.1	4F060568.1 NT		
Most Similar (Top) Hit BLAST E Veitue	1.0E-102 AF01	1.0E-102 AL163303.2	1.0E-102 BE252470.1	1.0E-102	1.0E-102 M10976.1	1.0E-102	1.0E-102	1.0E-102 BE40	1.0E-102 A1124	1.0E-102 A1124	1.0E-102	1.0E-102	1.0E-102 AU14	1.0E-102 AU1	1.0E-102 AL16	1.0E-102 BE2	1.0E-102	1.0E-103 BE90	1.0E-103 BE9(	1.0È-103 D87078.2	1.0E-103	1.0E-103 AJ27	1.0E-103	1.0E-103 AF0	1.0E-103	1.0E-103	1.0E-103	1.0E-103 AU1	1.0E-103 AFO	1.0E-103	1.0E-103 BE74472
Expression Signal	96.0	4.65	0.83	0.81	3.79	1.82	1.82	373.13	3.39	3.39	0.69	4.	2.88	2.88	1.63	2.09	1.17	1.86	1.86	7.11	6	0.87	8.76	3.06	0.99	1.28	1.28	2.16	1.58	1.28	2.58
ORF SEQ ID NO:	10108	10397	10645	10816	11137	11287	11288	11444	12342	12343	12824	13022	13089	13090	14091	14267	14903	10153	10154	10186	10284	11013	11269	11620	11847	12013	12014	12338		12630	
SEQ ID NO:	6120	5391	5642	5787	6107	6247	6247	6394	227	2227	7807	6008	8077	7208	9105	8278	9925		L		5272	5981	6223	6228	6828	6916	6916				Ш
Probe SEQ ID NO:	\$	339	616	882	138	1249	1249	1394	2246	2246	2786	2991	3060	3060	4111	4287	4948	88	88	68	208	996	1224	1561	1870	1930	1930	2242	2383	25.47	2995

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	Top Hit Descriptor	UI-H-BWO-GIRH-11-0-UI.st NCI_CGAP_Sub6 Hamo septens aDNA dans IMAGE:2733185 3'	Homo sepiens mRNA for KIAA1459 protein, perdiel cds	Macaca mulatta cyclophilin A mRNA, complete cds	ab (0d12 st Stratagene lung (#837210) Homo septems cDNA clone IMAGE:840407 3' similar to contains	element LTR10 repetitive element;	Homo sapiens neuropilin 1 (NRP1), mRNA	seq340 b4HB3MA-Cot109+10-Bio Homo sapiens cDNA clone b4HB3MA-Cot109+10-Bio-7 3	Hamo saplens chromosome 21 segment HS21C078	DKFZp564H1072_r1 664 (synonym: htbr2) Hamo septens CUNA data DN-72pown 1072 o	DKFZp564H1072_r1 564 (synonym: htbr2) Hamo sapiens CUNA date DN Zpockh 1072_r	Homo sapiens bone morphogenetic protein 8 (osteogenic protein 2) (pivir o) miruth	2022c06.s1 Stratagene colon (#83/204) Homo sepens cUNA done IMA/GE:08/020 3 Similar to gb:214116_ma1 CD59 GLYCOPROTEIN PRECURSOR (HUMAN);	601577460F1 NIH_MGC_9 Hamo saplens cDNA done IMAGE:3826438 o	RC1-CT0249-110900-214-f12 CT0249 Homo saplens cDNA	RC1-CT0249-110900-214-112 C10249 Homo sapiens cultiva	Homo saplens ARP2 (actin-related protein 2, yeast) nomoog (AC) rx2), mixton	Homo sapiens KIAA0440 protein (KIAA0440), mKNA	Homo saplens KIAA0440 protein (KIAA0440), mrnnA	Human lymphocytic antigen CD89/MEM43 mKNA, complete cus	H. Sapiens gene encouling prientypy uvale transmission in	AU133920 UVARCI nama seprens contra care of microscopic	ESTATIONS AGreened graine unition from 5 equations of 5 cm.	Trick A 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	HOUSTAND I DOMESTED THE RESIDENCE OF THE CONTROL OF	HUMAN MINIORICAN (TIN practiled)	Homo sapiens chromosome 21 unknown mistak	Homo sapiens chromosome 21 Unknown mrsky	Homo sepiens amyfold beta (A4) precursor protein (protease neadn-II, Abzheimer disease) (APP), mRNA	Homo sapiens Meis1 (mouse) homolog (MEIS1) mRNA	Homo expiens potassium channel subunit (HERG-3) mRNA, complaine cas	Homo saplens polassium channel subunit (HERG-3) mRNA, complete cds	Homo expiens mRNA for cyclin B2, complete cds	
2001	Top Hit Database Source	EST HUMAN				EST_HUMAN		EST_HUMAN		EST_HUMAN	EST_HUMAN		EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN						EST HUMAN	EST HUMAN	-	EST HUMAN			-	<u>-</u> -	L	1	E		
<b>}</b>		ES	E	į	Ļ	<u>8</u>	닏	ES	ż	ES	ES	FN 8	ES	ES	ES	ន	TONT	25 NT	<u>5</u>	Ē	될	띡	쁴	Ž	쁴	E	Į	토	<u>z</u> 8	1200	Z	본	Ž	1
Pilitino.	Top Hit Acesston No.	4W298245.1	4 OF 103 AR040892 4	4 0E-402 AE023881 4		AA485863.1	11430876 NT	T23683.1	AL163278.2	1.0E-104 AL037549.3	AL037549.3	4502428 NT	AA132976.1	BE744628.1	1.0E-104 BF334221.1	BF334221.1	5031570 NT	7682125 NT	7662125 NT	M34671.1	Y11151.1	1.0E-104 AU133928.1	1.0E-104 AA319436.1	1.0E-104 AB032998.1	1.0E-104 F11745.1	X02761.1	AF231920.1	AF231920.1	4502166 NT		4F03289	10		
	Most Stmiler (Top) Hit · BLAST E Vetue	1.0E-103 AV	4 OF-103	4 OF 403		1.0E-103 AV	1.0E-103	1.0E-103 T23683.1	1.0E-103	1.0E-104	1.0E-104	1.0E-104	1.0E-104	1.0E-104	1.0E-104	1.0E-104 BI	1.0E-104	1.0E-104	1.0E-104				1.0E-104	1.0E-104	1.0E-104	1.0E-104 >	1.0E-104	1.0E-104	1 0F-105			1.0E-105	L	
	Expression Signal	4.02	2	3 0	•	1.2	1.28	244	3.54	4.73	4.73	1.93	4.0	5.95	3.65	3.55	6.02	1.36	1.36		2.57	0.95	1.84	1.14	1.24	4.98	1.5		3 86					7.5
	ORF SEQ ID NO:	13331	100.70	19978		43714	18749	13895	14626		10306	11927				12403			12520	L		13227		13848	13992	14230	14448		40040		1			
	Exen SEQ ID NO:	8308	L	3 3	١	8708	8743	8897					l	L		L	L		L		7866	8205	8318	8841	9004			L				$\perp$	0700	╝
-	Probe SEQ ID NO:	3295		253	800	2704	2730	3897	4846	233	233	1849	2131	2141	2308	2308	2370	2430	2430	2800	2846	3189	3307	3839	4008	4252	4477	4477		8/7	1	200	8	86

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Top Hit Descriptor	Homo saplens chromosome 21 segment HS21C047	Homo saplens chromosome 21 segment HS21C080	Human mRNA for KIAA0128 gene, partial cds	EST20609 Solean I Homo sabiens CDNA 51 and similar to autoimmune extract K11 a zhinen autoimit	60143491F1 NIH MGC 72 Home saniens CDNA chine INA CE-2010511 K	no10d05.s1 NCI CGAP Phet Homo septems cDNA clone MAGE-11002653	Hamo sepiens 959 to carvia between AML1 and CBR1 on chromosome 21022, sepment 1/3	602022595F1 NCL CGAP Brn67 Homo sapiens cDNA clone IMAGE-4158143 F	602022595F1 NCI CGAP Brn67 Homo saniens cDNA clone IMA GF-4156143 ST	EST373761 MAGE resequences, MAGG Homo saplens cONA	Homo sapiens chromosome 21 segment HS21C008	Homo sapiens mRNA for KIAA0796 protein, partial cyls	Homo saplens mRNA for KIAA0866 protein, complete cds	EST378088 MAGE reseguences, MAGI Homo seriens china	UI-HF-BN0-akt-g-07-Q-UI-H NIH MGC 50 Homo saciens china clame MARCE: 3078348 FI	1q79c01.x1 NCI_CGAP_Ut1 Homo sapiens cDNA clone IMAGE:2215008.3	EST377629 MAGE resequences, MAGI Homo saciens cDNA	Human dihydrofolate reductase pseudogene (psi-hd1)	Human dinydrofolate reductase pseudogene (psi-hd1)	Homo sapiens soluble neuropilin-1 mRNA, complete cds	Human epidermal growth factor receptor (EGFR) precursor-mRNA, exon 4, partial cds	Homo sapiens type IV collagen alpha 5 chain (COL4A5) gene, exon 41	ng41-05.s1 NCI_CGAP_Co3 Homo sapiens cDNA clone IMAGE:937352.3' similar to contains element LTR3 repetitive element:	ng41c05.s1 NCI_CGAP_Co3 Homo sepiens cDNA clone (MAGE:837352.3' similar to contains element LTR3 repetitive element:	08-d10 HT0165 Homo seniens cDNA	AN	one (EDA), exon 2 and flanking repeat	regions	Homo saplens sparm membrane protein BS-63 mRNA, complete cds	601149783F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:3502461 5	q176h10.x1 Soares_NhHMPu_S1 Hamo saplens cDNA clone IMAGE:1878307 3"	Homo saplens glutathione S-transferase thela 1 (GSTT1), mRNA
Top Hit Deterbase Source	LZ LZ	NT TN	NT	EST HUMAN	EST HUMAN	EST HUMAN	N	EST HUMAN	EST HUMAN	EST_HUMAN	LN LN	N	N	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	LN LN	TN	Z,	N	NT	EST HUMAN	EST HUMAN	EST HUMAN	NT		NT L	NT	EST_HUMAN	EST HUMAN	Ę
Top Hit Acession No.	AL163247.2	AL163280.2	D50918.1	AA318369.1	BE891766.1	AA584808.1	AJ229041.1	BF347753.1	BF347753.1	AW961688.1	AL163208.2	AB018339.1	AB020673.1	AW966015.1	AW 503208.1	A1565065.1	AW965556.1	J00146.1	J00146.1	AF145712.1	U48724.1	U04510.1	AA527446.1	AA527446.1	BE144286.1	4504184 NT		+			A/276528.1	4504184 NT
Most Similar (Top) Hit BLAST E Value	1.0E-105	1.0E-105	1.0E-105	1.0E-105	1.0E-105		1.0E-105	1.0E-105	1.0E-105	_	_		1.0E-105	1.0E-105		1.0E-106	1.0E-106	1.0E-106	1.0E-106		_	1.0E-106	1.0E-106	1.0E-106	1.0E-106	1.0E-106		_				1.0E-106
Expression Signal	1.04	1.78	1.62	5.92	1.56	0.96	3.39	1.32	1.32	90.9	4.42	0.92	1.08	0.67	1.69	2.72	1.61	8.07	6.16	1.76	3.2	2.79	3.22	3.22	1.25	10.26		1.47	7	1.49	\$	7.19
ORF SEQ ID NO:	11738	11869		12220					13287	13964			14928	14938		10282	10576	10631	10631	11545	11732	11751	11845	11846	12153	12349		12522	12611	12613	12757	11462
Exan SEQ ID NO:	6663	6777	6874	7106	7237	7609	7956	8265	8265	8979	9765	9911	9950	9961	5216	5269	5572	5631	5631	6490	8999	6677	6229	6759	7045	7230		7403	7491	7493	784	6404
Probe SEQ ID NO:	1667	1785	1885	2126	2260	2649	2937	3252	3252	3981	4781	4934	4974	4987	150	202	537	602	603	1492	1662	1681	1767	1787	2063	2253		2432	2525	2527	2683	2752

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Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor		Hamo sapiens dutathione S-transferese thete 4 (DSTT4) DNA	801272675F1 NIH MGC 20 Home Control of the Control	Homo seriens mBNA for KIAA 4226	Homo seriens mRNA for Klas 41228 marking 12	Homo sapians throughout a national 144079 (T. 144079)	Homo seriens throughout antible 114273 (FLJ112/3), mKNA	Homo sapiens mBNA for Klas 4078 and 1.1.	Homo sapiens mRNA for KIA 41278 santis	Composations core blodies force alabel and an intervention of the composation force alabel and the	EST386875 MAGE resolutions MACAN Library	EST386875 MAGE resemblence, MAGN Lives seviers CONA	Home sarlons ADIS like 4 (ADIS 4) - DAIA	MRR-HT0165-140200 And 410 LT0455 L	Ilman alpha magneidocul II DANA	Figures applies mannestogase il mirava, comprete cas	inding separate Alphaeucodeucosomal region; segment 1/2	Human IFNAR gene for interferon alpha/beta receptor	numo seperis NT-KEN-25 anigen mKNA, pertiel cds	Human iFNAK gene for interferon alphafbeta receptor	Human IFNAR gene for interferon alpha/beta receptor	Homo sapiens sodium-decendent high-affinity dicarbox/date framenouter (NADCs) month	Homo sepiens BAZ18 mRNA for bromodomain adjacent to zinc finner domain 18 complete cas	QV2-HT0540-120900-358-e05 HT0540 Homo sapiens cDNA	Нато sapiens cathepsin Z precursor (CTSZ) gene, exon 3	Homo sapiens mRNA for KIAA0453 protein, partial cds	Homo saplens mRNA for KIAA0453 protein, partial cds	Human dipeptidy peptidase IV (CD26) gene, exon 20	PM1-CN0031-190100-001-d03 CN0031 Homo sapiens cDNA	PM1-CN0031-190100-001-d03 CN0031 Homo sapiens cDNA	601567619F1 NIH_MGC_21 Home septens cDNA clone IMAGE:3842309 5	601567619F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3842309 5	PM1-CN0031-190100-001-d03 CN0031 Homo sapiens cDNA	PM1-CN0031-190100-001-d03 CN0031 Homo sapiens cDNA	Homo sapiens SMT3 (suppressor of mif two 3, yeast) homolog 2 (SMT3H2), mRNA
	Top Hit Detabase		L <sub>V</sub>	EST HUMAN	LZ	LZ	l'a	15	NT.	Į	TN	EST HUMAN	EST HUMAN		EST HIMAN	L		<u> </u>	Ž	-	- N	LN.	_ <del>\</del>	F	EST_HUMAN	TN	NT		TN	EST_HUMAN	П				T_HUMAN	
	Top Hit Acession No.		4504184 NT	E384296.1	B037747.1	B037747.1	8922965	8922965 NT	1.0E-106 AB033104.1					9226		Γ	T	T	1				154121.1						13729.1	N842451.1						5902097 NT
10.00	(Top) Hit BLAST E	Value	1.0E-106	1.0E-106 B	1.0E-106 A	1.0E-106	1.0E-106	1.0E-106	1.0E-106	1.0E-108	1.0E-106	1.0E-106 A	1.0E-106 A	1.0E-106	1.0E-106 BE144286.1	1.0E-106 U31520.1	1.0E-107 A.1274735 4	4 OE 407 VENAED 4	1.0E-107 AE4EE409 4	4 OF 407 X	1.0E-107 A00439.1	7.0E-10.7	1.0E-107 AF	1.0E-107 AB032253.1	1.0E-107 BF087405.1	1.0E-107 AI	1.0E-107 AB007922.2	1.0E-107 AB007922.2	1.0E-107 U	1.0E-107 A	1.0E-107 A	1.0E-107 BE732460.1	1.0E-107 BE732460.1	1.0E-107 AW842451.1	1.0E-107 AW842451.1	1.0E-107
	Expression Signal		7.19	1.79	4.45	4.45	2.41	2.41	0.99	66.0	0.92	10.11	10.11	1.74	0.67	1.39	3.3	4 47	2 2 2	104	10.1	C7.L	8.1	2.22	8.86	4.61	2.87	2.87	1.06	4.22	4.22	1.65	1.65	3.71	3.71	7.53
	ORF SEQ ID NO:			12840	12910		13141	13142	13380	13381	13719	13916	13917	13932	14439	14711			10655	10854	7000	+080	11003	11300	11593	11784	11885	11886	12242	12395	12396	12550	12551	12974	12975	13052
	SEQ ID						8123	8123			8718	8926	8926	8942	9460	9726	5297	5321	5651	5824	5803	200	5970	6257	6534	6708	6795	6795	7125	7276	7276	7432	7432	7957	7957	8043
	Probe SEQ ID NO:		2752	2805	2870	2870	3107	3107	3356	3356	3714	3926	3926	3944	4470	4741	234	262	624	803	R75		954	1259	1536	1713	1804	1804	2146	2301	2301	2483	2463	2938	2938	3026

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Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo sapiens metubulerin (MTM1) cane arm o	601177018F1 NIH MGC 17 Home canleng c ONA chara MAA CE 32372 49 21	Homo sapiens NF2 cene	601671914F1 NIH MGC 20 Homo seniens cDNA close NAACE 3854030 E'	bb25b10.x1 NIH_MGC_14 Homo saplens cDNA clone fMAGE:2963699 3' similar to gb:X53777 60S RIBOSOMAL PROTFIN 1.29 (HI MAAN): nb:2727 Marine homelessessessesses)	M12a11x1 NG_CGAP_GUT Home septems cDNA clone IMAGE:2972080 3' similar to SW:38P1_MOUSE P55194 SH3-BINDING PROTEIN 3RP-1 .	Human hepatocyte nuclear factor 4-alpha gene axon 2	Human hepatocyte nuclear factor 4-alpha cene avon 2	Homo sapiens KIAA0187 gene product (KIAA0187) mRNA	UI-HF-BNO-ain-e-04-0-UI.r1 NIH MGC 50 Home sapiens china MAAGE 3080168 F	Homo saplens PSN1 gene, alternative transcript	Homo sapiens CCAAT-box-binding transcription factor (CBF2) mRNA	Homo saplens mRNA for Golgi-associated microtubule-binding protein (GMAP-210)	IL2-UM0077-260400-079-D06 UM0077 Homo septens cDNA	Human mRNA for KIAA0220 gene, partial cds	Homo saplens hypothetical protein FLJ11316 (FLJ11316), mRNA	Homo sapiens reticulocalbin 1, EF-hand calcium binding domain (RCN1), mRNA	Homo sapiens tetratricopeptide repeat domain 2 (TTC2) mRNA	Homo sapiens mRNA for KIAA0999 protein, partial cds	Homo sapiens mRNA for KIAA0999 protein, partial cds	Homo sapiens nucleolar phosphoprotein B23 (NPM1) mRNA, complete cds	Homo sapiens nucleolar phosphoprotein B23 (NPM1) mRNA, complete cds	601186922F2 NIH_MGC_15 Hamo sapiens cDNA clone IMAGE:2859636 5	601186922F2 NIH_MGC_15 Hamo sepiens cDNA clone IMAGE:2959636 5	Homo saplens mRNA for KIAA0018 protein, partial cds	Homo sapiens chromosome 21 segment HS21C084	Homo sepiens SNF&INI1 gene, exon 6	ow95e01.x1 Soeres_fetal_liver_spleen_1NFLS_S1 Höme saplens cDNA clone IMAGE:1654639 3' similar to TR:002197 002197 CRCULATING CATHODIC ANTIGEN.;	ow95e01.x1 Scares_feta_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:1654536 3' similar to TR:002197 002197 CIRCULATING CATHODIC ANTIGEN;
Second Lines	Top Hit Database Source	L'A	EST HUMAN	N P	EST HUMAN	EST HUMAN	EST HUMAN	Į. Į.	TN.	F	EST HUMAN	N	<u> </u>	LN	EST_HUMAN	NT	<del>را</del>	47	47	NT	NT	NT	NT	EST_HUMAN	ST_HUMAN	NT	NT	NT	EST_HUMAN	EST_HUMAN
1 916 110	Top Hit Acession No.	AF020671.1	BE296042.1		BF026728.1	BE206694.1	Í _	U72961.1		7661979 NT	1.0E-108 AW504799.1	1.0E-108 AJ008005.1	5031624 NT		1.0E-109 AW803116.1		11422486 NT	11438391 NT	7712		-				Ψ.		1.2		1022328.1	1.0E-109 A1022328.1 E
	Most Similar (Top) Hit BLAST E Value	1.0E-107	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108	1.0E-108 Y12490.1	1.0E-109	1.0E-109 D86974.1	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109	1.0E-109 M28699.1	1.0E-109 M28699.1	1.0E-109	1.0E-109	1.0E-109 D13643.2	1.0E-109	1.0E-109 Y17123.1	1.0E-109 A	1.0E-109
	Expression Signal	5.33	2.88	5.83	1.65	12.25	1.15	3.04	3.04	3.68	0.88	2.91	0.95	0.72	2.46	3.94	0.76	7.69	9.2	15.28	15.28	77.88	72.21	1.52	1.52	5.54	2.32	4.31	4.09	4.09
	ORF SEQ ID NO:	13739	10990	11284	12114	12451	14017	14367	14368	14632	14737											-				11914	12280	12288	12632	12633
	Exan SEQ ID NO:	l			7008	7334	9029	9385	9385	9644	9749	9775	8939	9962	5123	5145	5281	5288	5501	5624	5624	6185	6185	6508	6508	6826	718	7168	7514	7514
	Probe SEQ ID NO:	3736	8	1246	2025	2360	4033	4394	4394	4659	4765	4791	4962	4988	43	99	218	922	464	293	293	1183	1184	1510	1510	1836	2181	7 <u>8</u>	2549	2549

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Single Exon Probes Expressed in HBL100 Cells

nu93c12.s1 NCI\_CGAP\_Pr22 Homo sepiens cDNA clone IMAGE:12182623' similar to SW:GTT2\_HUMAN nu93c12.s1 NG\_CGAP\_P722 Hamo sepiens cDNA done IMAGE:12182623' similar to SW:GTT2\_HUMAN Homo sapiens Bruton's tyrosine kinase (BTK), apha-D-galectosidase A (GLA), L44-like ribosomal protein 12816F Human fetal heart, Lambda ZAP Express Home sephens cDNA clone J2816 5 strutter to ZINC ts98e06.x1 NCI\_CGAP\_GC6 Home sapiens cDNA clone IMAGE:2239330 3' simitar to WP:F53A2.8 Homo sapiens BAZ18 mRNA for bromodomein adjacent to zinc finger domain 18, complete cds 601237545F1 NIH\_MGC\_44 Homo sapiens cDNA clone IMAGE:3609683 5 Homo sapiens BAZ1B mRNA for bromodomain adjacent to zinc finger domain 1B, complete cds 0.032b/0.x1 Sogres\_NFL\_T\_GBC\_S1 Hamo sapiens cDNA clane IMAGE:1627863 3' similar to SW:N121\_RAT P52591 NUCLEAR ENVELOPE PORE MEMBRANE PROTEIN POM 121; UI-H-BI4-acs-b-05-0-UI.s1 NCI\_CGAP\_Sub8 Home sepiens cDNA clone IMAGE:3085784 3 Homo saplens chondratin sulfate proteoglycan 4 (melanoma-associated) (CSPG4), mRNA Homo sapiens retinol dehydrogenase homdog ladform-1 (RDH) mRNA, complete cds MR0-HT0209-110400-108-s04 HT0209 Homo sapiens. cDNA Homo sepiens deiodinase, iodothyronine, type II (DIO2), transcript variant 2, mRNA Homo sapiens deiodinase, todothyronine, type II (DIO2), transcript variant 2, mRNA Homo sapiens leucine zipper-like transcriptional regulator, 1 (LZTR1), mRN/ Homo sapiens leucine zipper-like transcriptional regulator, 1 (LZTR1), mRN/ ruman autoimmune antigen small ruclear ribonucleoprotein E pseudogene Homo sapiens guanylate cyclase activator 1A (retina) (GUCA1A) mRNA Homo sapiens KIAA0377 gene product (KIAA0377), mRNA Homo sapiens guanylate cyclase activator 1A (retina) (GUCA1A) mRNA AU117812 HEMBA1 Homo sapiens cDNA done HEMBA1002241 5" Human mRNA for inward rectifier potassium channel, complete cds Top Hit Descriptor CM3-NN0009-190400-150-f10 NN0009 Homo septens cDNA CM3-NN0009-190400-150-110 NN0009 Homo septens cDNA P30712 GLUTHATHIONE S-TRANSFERASE THETA 2; P30712 GLUTHATHIONE S-TRANSFERASE THETA 2; Homo sapiens calcitonin receptor-like (CALCRL) mRNA Homo sepiens KIAA1002 protein (KIAA1002), mRNA (L44L) and FTP3 (FTP3) genes, complete cds Human dystrobrevin (DTN) gene, exan 20 FINGER PROTEIN ZNF43 CE16100 EST HUMAN EST\_HUMAN HUMAN HUMAN Top Hit Database Source EST HUMAN EST\_HUMAN EST\_HUMAN EST\_HUMAN EST\_HUMAN EST\_HUMAN EST HUMAN EST **EST** 5803073 NT 5803073 NT þ 4504206 NT 7662083 NT 7549804 NT 4504206 NT E 눋 눋 7549804 NT 5031620 NT 4503098|NT Top Hit Acessian 7662441 1.0E-109 AW 893 192, 1 1.0E-109 AA662274.1 AW893192. 1.0E-109 BE146144.1 1.0E-109 AA662274.1 AB032253.1 AF240698.1 ġ 1.0E-109 AIB55417.1 I.0E-110 BE379477.1 1.0E-110 AU117812.1 1.0E-110 AB032253.1 1.0E-110 AI017213.1 1.0E-109 N85190.1 U84550.1 M15918.1 1.0E-110 U78027.1 1.0E-110 ( 1.0E-109 1.0E-109 1.0E-110 1.0E-109 .0E-110 1.0E-110 1.0E-110 1.0E-110 (Top) Hit BLAST E 1.0E-109 1.0E-110 1.0E-110 1.0E-110 1.0E-110 Most Simila 1.32 43 4.42 1.43 2.78 0.94 209 241 1.07 3.85 0.79 0.83 0.78 0.8 1.29 1.07 2.3 0.92 2.55 3.28 Expression 13014 13342 14006 14022 14262 14438 10068 9106 ORF SEQ 12634 10105 0068 10362 11199 14062 14466 10561 11301 11957 11301 14494 347 ÖZ 8002 8315 8315 8448 8755 9019 5084 5118 SEQ ID 7515 9034 9034 9459 5118 5349 6868 9510 6165 9798 6988 9074 9490 6258 6258 88 2 2550 30 30 20 SEQ ID 3440 3751 \$ 5 28 28 28 28 28 28 2005 5023 1038 292 523 523 169 8 8 5 1260 4500 4520 4814 2961 3013 | <del>6</del>

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Human ribosomal protein L23e mRNA, complete cds	Homo sapiens ras GTPase activating protein-like (NGAP) mRNA	GNASSES1F1 NIH MGC 68 Homo saplens oDNA clone IMAGE:3862086 5	User Control on the surface critical region come 1 (CECR1). mRNA	Figure September 19 89 Interior before the first of the second 32 to 34	HUMBIN CARRIED STATES CHAIR (WILL OF BACK)	Homo sepiens Nikkupup gerie julianak (Nikezbazanaka) mana	Homo septents UNITER4340 (30 proven) (UNITER 1530) (11 provens Camplete cds	Human enkepriami o (emo) gene, emis e mare en emis emis en emis emis emis en emis emis emis en emis emis emis emis emis emis emis emis	Homo sepiens acetyl-coencynters cellularyiass bota (notae sign)	Human sterodogenic acute regulatory procein (John's) gone, come	Human sterodogenic acute regulatory protein (2011) years, waste of the live of the land live of the li	UI-H-BI4-BI4-BI4-UI-SI NOT COMP CHICA CAPACITO COMPANIA CHICA CAPACITO COMPANIA CHICA CHICA CAPACITO CHICA C	UI-H-BI4-B04-0-U.ST NCI COAT Suito nuito advicto con constante cos	Homo sapiens H IKA same protesse (Truch II) yard, vorignos de	ZING FINGER PROTEIN 135	Homo sapiens KIAA0440 protein (KIAA0440), mKNA	Homo saplens KIAA0440 protein (KIAA044u), mixink	601442674F1 NIH_MGC_65 Homo sapiens CUNA done IMAGE: 3040809 9	Homo sapiens glutamate receptor, tonotropic, Kainate 1 (GKIN1) mixivA	RC2-BT0642-030400-021-d09 BT0642 Homo sepiens cDNA	RC2-BT0642-030400-021-409 BT0642 Homo sapiens cDNA	MR2-BT0590-090300-113-f09 BT0590 Homo sapiens cDNA	Homo sapiens mRNA for KIAA1411 protein, partial cds	Homo sapiens mikina tor Kida 141 potent, parter cus	acgorot, x1 Schiller meningiona from septems contraction in the contra	ao95f01,x1 Schiller meningionna riomo saprans con numbra:	Human X-linked phosphoglycerate kinase gene, exert o	ao95f01x1 Schiller meningioma Homo sapiens curva cione invoca.	Homo sapiens elf4E-transporter mRNA, complete cds	Homo sapiens mRNA for putative RNA helicase, 3 end	Homo sapiens mRNA for multidrug resistance protein 3 (ABCC3)	Homo sapiens mRNA for multidrug resistance protein 3 (ABCC3)	Homo sapiens mRNA for multidrug resistance protein 3 (ABCC3)	
Top Hit Database Source			T LI BAAN	NO.									EST HUMAN	EST HUMAN		SWISSPROT		-	EST_HUMAN		EST HUMAN	EST HUMAN	EST HUMAN			EST HUMAN	EST HUMAN	<b>_</b>	EST HUMAN	Z.	Į.	Į	Ę	12	
	눌	Į,			Z I	칠	취	되	뵈	<u> </u>	Ż	Ż	ည္	篮	호	S	25 NT	IN 52	낊	<u>S</u>	li i	T W	Ä	N	E	Ü	E	ΤN	Ü	Z	F	Z	+	12	
Top Hit Acession No.	143701.1	A7E0BO7	4/3000	1.0E-111 BF035327.1	8393092 NT	5142	7662177 NT	7661569 NT	K02268.1	4501854 NT	J29103.1	U29103.1	1.0E-112 BF509039.1	1.0E-112 BF509039.1	1.0E-112 AF157623.1	P52742	7662125 NT	7662125 NT	BE86685	4504116 NT	4 AE 442 BEARANG2 1	4 0F-112 BE083092.1	1 0F-112 BE076073.1	1.0E-112 AB037832.1	1.0E-112 AB037832.1	1.0E-113 Al365586.1	1.0E-113 AI365586.1	M11965.1	Al365586.1					2.161717	
Most Similar (Top) Hit BLAST E Value	1.0E-111 U43701.1	100.	1.0E-111	1.0E-111	1.0E-111	1.0E-111 M2	1.0E-111	1.0E-111	1.0E-111	1.0E-112	1.0E-112 U29103.1	1.0E-112 U29103.1	1.0E-112	1.0E-112	1.0E-112	1.0E-112 P52742	1.0E-112	1.0E-112	1.0E-112	1 0F-112	4 OF 142	4 0F-112	1 OF-112	1.0E-112	1.0E-112	1.0E-113	1.0E-113	1.0E-113	1 0E-113			ł	١		1.0E-114
Expression Signal	43.9		1.07	. 2.71	3.66	2.29	1.57	1.17	4.64	0.87	4.82	4.82	1.33	1.33	1.84	2.53	3.11	3.11	1 26	52.0	200	1.07													0.76
ORF SEQ ID NO:	+	1	10273		10779	10965	11648	14032	14178	10632	10633					l						13208	1	1	1.										10140
Exon SEQ ID NO:	603	9530	5260	5748	5757	5930	6586	9043	9196	5632				5653	6001	L			1	١			┙	9595		L	1								5139
Probe SEQ ID NO:	1	7	196	725	734	914	1589	4047	4203	909	909	909	626	626	OBG	1045	1045		2000	2430	3004	3171	3171	3/90	8087	2 2	3 5	3 8	Ä	1509	1899	3057	ຜິ	29	Ğ

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Table 4
Single Exon Probes Expressed in HBL100 Cells

																_		_		_	_	<u> </u>	"Lul		~	~	-	$\overline{}$	_				
	Top Hit Descriptor	yd15c01.st Soares four liver spleen 1NFLS Homo septens cDNA clone IMAGE:108288 3' shriifar to gb:A21187 ALP4IA-2AAGROGLOBULIN PRECURSOR (HUMAN);contains Alu repetitive element;	Homo sapiens hypothetical protein TLLZQUed (TLLZQUed), ninver	Hamp septents metadata wind defendi inglikat protein (1000-100) (1000-100) mRNA	Homo sapiens minicipromosome memierianos delicien (5. cerenistas) o (morro), minicipalita (5. cerenistas) o (morro), minicipal	Homo sapiens ruckeoporin-like protein 1 (NLP 1), miruth	Human mKNA 10' KIAAU3/0 gene, parua cos	Homo sapiens miking to NIAA 1270 propert, per use was	Home sapiens miking for NIAA1270 protein, parter to chemicana 41 hand n/3	Human gene for catalase (E.C. 1.11.1.6) aron 2 mapping to children in your price.	601869932F1 NIH MGC 19 Homo sapiens CANA clore invacing to the control of the con	Homo saplens NOD1 protein (NOD1) gene, exons 1, 2, and 3	Human interferon-alpha receptor (Huit-N-alpha-Kec) mrunh, complete cus	601122173F1 NIH_MGC_20 Hamo sapiens cDNA clone IMAGE:3346099 5	Homo sapiens HLA-B associated transcript-1 (D6S81E) mRNA	Homo sapiens polymerase (RNA) II (DNA directed) polypeptide A (220kD) (POLK2A) mKNA	Homo saplens keratin 18 (KRT18) mRNA	QV4-UNX094-300300-156-b08 UM0094 Homo saplens cDNA	Homo sapiens transforming growth factor beta-activated kinase-binding protein 1 (TAB1), mRNA	Homo saplens transforming growth factor beta-activated kinase-binding protein 1 (1A51), mixiva	Homo sapiens ferritin, heavy polypeptide 1 (FTH1) mtKNA	Homo sapiens alpha-aminoadipate semialdehyde synthase mitNA, complete cas	Homo saplens alpha-aminoadipate semialdenyde symmase mr.N.A. compress cus	Homo sapiens Bruton's tyrosine kinase (BTK), apha-U-galectosidase A (CLA), L44-lina incomina promin (L44L) and FTP3 (FTP3) genes, complete cds	601579838F1 NIH_MGC_9 Homo saplens cDNA clone IMAGE:3928832 5	601579838F1 NIH MGC 9 Homo sepiens cDNA clone IMAGE:3928832 5	Homo saplens testican-1 mRNA, complete cds	10/4 HA0094-300300-156-b08 UM0094 Homo sepiens cDNA	Lorens englans mRNA for alpha tubulin 8 (TUBA8 gene)	Trong seriors mRNA for alpha-trbulin 8 (TUBA8 gene)	Tions soules notice I'm done for tith	Home society mRNA for KIAA0350 protein, partial cds	
	Top Hit Database Source	T_HUMAN								- 1	ST_HUMAN	_	l-	EST HUMAN	1		_	EST HUMAN	1		F	Ţ	<b>J</b>	<u>_</u>	EST HUMAN	EST HIMAN	F17	ET CHIMAN	NUMBER 181	Ž!	Z	Z	=
2	ion	EST	8923087 NT	7657529 NT	6631094 NT	6679073 NT	Ż	Z	Z	Z	EST	TN	Z	S	4758111 NT	4505938 NT	4557887 NT	Γ	4702	5174702 NT	4503794 NT	LN	Z	눌	Ü		)  2	T	1				7
2	Top Hit Acession No.	170551.1	892	765	663	6679	AB002374.1	AB033102.1	AB033102.1	X04086.1	BF206374.1	AF149773.1	J03171.1	BE275324.1	475	450	. 455	AW804759.1	517			AF22918	AF229180.1	178027.1							AJ245922.1		AB002348.2
	Most Similar (Top) Hit BLAST E Value	1.0E-114	1.0E-114	1.0E-114	1.0E-114	1.0E-114	1.0E-114	1.0E-114	1.0E-114	1.0E-114	1.0E-114	1.0E-114	1.0E-114	1.0E-114	1.0E-115	1 0E-115	1.0E-115	1 0E-115	1 0E-115	1.0E-115	1.0E-115	1.0E-115	1.0E-115	1 05-115	4 OF 445	400 445	1.0E-113	1		_	١	-	1.0E-115
	Expression Signal	5.07	3.1	3.85	5.21	11.15	1.31	1.1	1.1	2.52	1.9	1.56	1.12	1.66	11.8	27.0	21.76	4 40	4.42	1 42	83.62	1.41	1.41	000			1.38	3.07	2.22	7.57			5.04
	ORF SEQ ID NO:	10869	11093	11338	11659	11699	12294	10114	10115		13132	13905	]_						1030F				_					12330		13067	13068		13920
	Exan SEQ ID NO:	5885	8083	6292	6598	9630	7173	5125	5125	8075	8114	8909	9259	9941	5102	100	5200	200	5200	5700	5804	L	L	1		210/	7010	ı	7799	8028	8028		8929
	Probe SEQ ID (9 NO:	83	1054	1294	1602	1633	2194	2732	2732	3058	3008	3008	4266	750A	3 6	1 5	25	5 5	250	9 6	2 00	4528	4528	2	1805	2027	2027	2236	2778	3041	3041	3392	3929

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Human offactory receptor olfr17-201-1 (OR17-201-1) gene, offactory receptor olfr17-32 (OR17-32) gene and Human apolipoprotein B-100 (apoB) gene, exons 17 and 18 Homo sapiens protein phosphatase, EF hand calcium-binding domain 1 (PPEF1) mRNA 4932c11.s1 Soares\_NFL\_T\_GBC\_S1 Homo sapiens cDNA clone IMAGE:1578548 3 Novel human mRNA from chromosome 1, which has similarities to BAT2 genes Novel human mRNA from chromosome 1, which has similarities to BAT2 genes 601121347F1 NIH\_MGC\_20 Homo septens cDNA clone IMAGE:2988875 5 Mus musculus fragile-X-related protein 1 (Fxr1h) gene, exons 13a through 15 Homo sapiens lymphocyte activation-associated protein mRNA, complete cds 601513337F1 NIH\_MGC\_71 Homo saplens cDNA clone IMAGE:3914600 5 olfactory receptor pseudo\_olfr17-01 (OR17-01) pseudogene, complete cds Homo sepiens acety-Coenzyme A carboxylase alpha (ACACA), mRNA AU133080 NT2RP4 Homo sapiens aDNA done NT2RP4001228 5 Hamo sapiens DiGeorge syndrome critical region, centrameric end Homo sapiens DiGeorge syndrome critical region, centromeric end Homo sapiens hypothetical protein FLJ10466 (FLJ10466), mRNA Homo sapiens hypothetical protein FLJ10466 (FLJ10466), mRNA Top Hit Descriptor Homo sapiens sodium phosphate transporter 3 (NPT3) mRNA Human apolipoprotein B-100 (apoB) gene, exon 10 EST369769 MAGE resequences, MAGE Homo saplens cDNA Human apolipoprotein B-100 (apoB) gene, exons 17 and 18 Homo saplens partial 5-НТ4 receptor gene, exons 2 to 5 Homo sapiens mRNA for KIAA0790 protein, partial ods Homo sapiens chromosome 21 segment HS21C068 PM-BT135-070499-016 BT135 Homo sapiens cDNA Homo sapiens chromosome 21 segment HS21C068 -fomo sapiens synaptojanin 1 (SYNJ1), mRNA Human alpha-5 collagen type IV gene, exon 5 Novel human gene mapping to chomosome X Homo sepiens sir2-like 3 (SIRT3), mRNA Homo sapiens pericentrin (PCNT) mRNA Homo sapiens pericentrin (PCNT) mRNA Homo sapiens EphA4 (EPHA4) mRNA Single Exon Probes Expressed in HBL100 Cells BINT EST\_HUMAN NT EST HUMAN Top Hit Database Source HUMAN EST HUMAN EST 6912659|NT 5453941 NT 눋 F 8922435 NT 5174478 NT 눌 5031954 NT 8922435 Top Hit Acession 4507334 5174478 4826636 .0E-116 AU133080.1 1.0E-115 AL137163.1 AL163268.2 AL096857.1 1.0E-115 AL096857.1 .0E-116 AJ243213.1 BE889256. AF123320.1 1.0E-116 AB018333. 1.0E-117 AA978114 1.0E-116 M19824.1 .0E-116 M19824.1 1.0E-116 U78308.1 1.0E-117 M19816.1 1.0E-117 M63468.1 1.0E-116 L77570.1 .0E-115 1.0E-115 1.0E-115 1.0E-115 1.0E-116 .0E-115 1.0E-116 (Top) Hit BLAST E 1.0E-115 1.0E-116 .0E-116 1.0E-116 1.0E-116 1.0E-117 fost Similar 1.0E-117 .0E-1 3.78 0.91 2.4 2.89 1.39 Expression Signal 0.91 2.12 38 86. 0.95 206 1.27 7 3.05 <del>ا</del> 1.93 ORF SEQ ID NO: 14515 14516 14429 14289 14430 14721 10599 10843 12038 12039 12066 12480 13135 14675 14945 10589 1785 11876 12245 12621 12131 13232 9139 9271 9449 9305 SEO ID 9449 9529 9529 9735 9735 5813 6937 7255 7710 8117 8117 9247 9692 5588 6937 0969 4313 4144 4459 4539 4539 4539 792 4750 2279 2387 2660 3101 SEQ ID 1975 2040 2040 3101 567 1253 951 554 4707 1061

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	EST188414 HCC cell line (matastasis to liver in mouse) II Homo sepiens cDNA 5' end similar to ribosomal protein L29	Homo saciens collegen have N alpha 5 (Albort explorate) (CO) 44.51 Chick	DKFZ0434C1120 rt 434 (strong m: Hecs) U	H. Sapiens mRNA for TPCR46 modeln	H. Sapiens mRNA for TDCR16 motein	Homo sepiens Scar (SCAR2) none partial cuts	Homo sapiens Scar 2 (SCAR2) dene mental cuts	Homo saplens mRNA for KIAA0868 mindeln complete cute	Homo septens HSPC151 mRNA, complete ods	DKFZp4341056 r1 434 (synonym: https://www.saniens.chwa.chwa.chwa.chwa.chwa.chwa.chwa.chwa	Homo saplens hypothetical protein (D.1328F19 C1.11 mPNA	Homo sepiens sine oculis homeobax (Drosophila) homolog 1 (SIX1) mRNA	601281947F1 NIH MGC 44 Homo sapiens cDNA clone IMAGE: 3674016 F	601281947F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3604019 5	601281947F1 NIH_MGC_44 Hamo sepiens cDNA clone IMAGE:3604019 5	ES1363799 MAGE resequences, MAGB Homo sapiens cDNA	Himan breathnaid cliebs sector (BCR) gene, complete cds	Homo sapiens PRKY exon 7	ap01f05.x1 NCI_CGAP_Kid5 Homo septens cDNA clone IMAGE:1916769 31	qp01f05.x1 NCI_CGAP_Kid5 Homo sapiens cDNA done MAGE:1916769 3"	Human mRNA for ribosomal protein, complete cds	Homo sapiens KIAA0478 gene product (KIAA0478), mRNA	Homo sapiens chloride channel CLC4 (CIC4) mRNA, complete cds	Homo saplens CGI-105 protein (LOC51011), mRNA	Homo sapiens mRNA for KIAA0930 protein, partial ods	Homo sapiens hypothetical protein FLJ10052 (FLJ10052), mRNA	Homo sapiens glutamete receptor, Ionotropic, kainate 1 (GRIK1) mRNA	Homo saplens synaptojanin 1 (SYNJ1), mRNA	Homo sapiens Intersectin 2 (SH3D1B) mRNA, complete cds	Homo saplens intersectin 2 (SH3D1B) mRNA, complete cds	1940g12.r1 Soares melanocyte 2NbHM Homo sapiens cDNA clone IMAGE:273769 51
Top Hit Database Source	MAN		MAN				Ī				Ī		Г		П	丁	Ī	Ī	Γ	П	-	1		_	•	1		+			
. 6 "	EST HUMAN	L	EST HUMAN	Į	z	上	Ę	Ę	E	EST_HU	Ę	¥	ST HUN	EST_HU	STH			  -	ST_HU	ST HU	Ϋ́	F	Ļ	늘	F	17	17	T.	5	ş	ST_HI
Top Hit Acession D	A316723.1	39564	T	T	89670.1				Γ	AL045854.1 EST_HUMAN	7657016 NT	5174680 NT		BE389705.1 EST_HUMAN	E389705.1	W851729.1 EST HUMAN		3932.1	A1347694.1 EST_HUMAN	EST	723660.1 NT	11425783 N.T	17049	5607	AB023147.1 N.T	8922205 NT	4504116 NT	4507334 NT		1.1	44873.1 EST_HUMAN
	1.0E-117 AA316723.1	39564	1.0E-117 AL042120.1 EST HI	T		F134304.2		1.0E-117 AB020673.1 NT		AL045854.1	1.0E-118 7657016 NT				E389705.1	T		13932.1	1.0E-118 Al347694.1 EST_HU	1 EST	D23660.1		AF170492.1	5607	AB023147.1			1.0E-120 4507334 NT		1.1	1.0E-120 N44873.1 EST_HI
Top Hit Acession No.	A316723.1	8659564	T	T	89670.1				Γ	1.0E-118 AL045854.1		1.0E-118	1.0E-118 BE389705.1	1.0E-118 BE389705.1	1.0E-118 BE389705.1	5	1.0E-118 U07000 1	1.0E-118 Y13932.1		1.0E-118 Al347694.1 EST	1.0E-118 D23660.1	1.0E-118	1.0E-119 AF170492.1	1.0E-119 7705607	1.0E-119 AB023147.1	1.0E-119	1.0E-119	1.0E-120	1.0E-120 AF248540.1	1.0E-120 AF248540.1	
Most Smiler (Top) Hit Top Hit Acession BLAST E No.	13885 14.06 1.0E-117 AA316723.1	1.0E-117 8659564	1.0E-117 AL042120.1	1.0E-117 X89670.1	1.0E-117 X89670.1	1.0E-117 AF134304.2	11.6 1.0E-117 AF134304.2	3.85 1.0E-117 AB020673.1	11.76 1.0E-118 AF161500.1	1.94 1.0E-118 AL045854.1	7.24 1.0E-118	6.59 1.0E-118	7.75 1.0E-118 BE389705.1	7.75 1.0E-118 BE389705.1	1.0E-118 BE389705.1	1.0E-118 AW 951 729.1	3.94 1.05-118 (107000 1	4.44 1.0E-118 Y13932.1	4.93 1.0E-118 Al347694.1	4.93 1.0E-118 Al347694.1 EST	17.63 1.0E-118 D23660.1	1.42 1.0E-118	0.93 1.0E-119 AF170492.1	4.3 1.0E-119 7705607	3.42 1.0E-119 AB023147.1	0.92 1.0E-119	0.86 1.0E-119	1.07 1.0E-120	2.23 1.0E-120 AF248540.1	2.23 1.0E-120 AF248540.1	1.0E-120 N44873.1
Most Similar  Expression (Top) Hit Top Hit Acession Signal BLAST E No.	13885 14.06 1.0E-117 AA316723.1	2.65 1.0E-117 8659564	14417 2.36 1.0E-117 AL042120.1	14561 1.11 1.0E-117 X89670.1	1.11 1.0E-117 X89670.1	14638 11.6 1.0E-117 AF134304.2	14639 11.6 1.0E-117 AF134304.2	14750 3.85 1.0E-117 AB020673.1	10155 11.76 1.0E-118 AF161500.1	10181 1.94 1.0E-118 AL045854.1	10553 7.24 1.0E-118	10957 6.59 1.0E-118	12267 7.75 1.0E-118 BE389705.1	12268 7.75 1.0E-118 BE389705.1	7.75 1.0E-118 BE389705.1	12738 3 94 1 0E-118 AW 951729.1	12739 3.94 1.0E-118 U07000 1	4.44 1.0E-118 Y13932.1	13159 4.93 1.0E-118 Al347694.1	13160 4.93 1.0E-118 Al347694.1 EST	13958 17.63 1.0E-118 D23660.1	14545 1.42 1.0E-118	10797 0.93 1.0E-119 AF170492.1	11059 4.3 1.0E-119 7705607	11972 3.42 1.0E-119 AB023147.1	13056 0.92 1.0E-119	13857 0.86 1.0E-119	10368 1.07 1.0E-120	11065 2.23 1.0E-120 AF248540.1	11066 2.23 1.0E-120 AF248540.1	8.07 1.0E-120 N44873.1

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Homo sanions deinformin and motollameters	Homo contact activities in the same of the	Homo services gains for AE-8 commissionals	Homo septens superplants 4 (SVA 14) — Data	Hampening CAMB - config the thinking	Homo series calific phosphodischess, 84 (FUEGA) mitha, partial cds	Homo saviens standardein /CTC/ and and all	Homo socione standardin (510) gara; partial 11	Homo sapians NE2 years	AU134963 PLACE1 Homo seriens CDNA when BLACE4000000 El	Hamp sepiens TNF reportitional factor of TPAEAT COMIA	602014759F1 NCI CCAP Bried Home service 2014	602014759F1 NCI CGAP Brink Home seniors cDNA close into Act 4420286 5	Homo sepiens serties palmithout transferese entrantil social and a sepiens serties palmithout transferese entrantil social and a sepiens serties palmithout transferese entrantil social and a sepiens serties palmithout transferese entrantil social and a sepiens serties palmithout transferese entrantil social and a sepiens serties	Homo sapiens hHb3 gene for hair keratin exms 1 to 0	Homo sapiens hHb3 gene for hair keratin axons 1 to 0	Homo saplens mRNA for KIA41337 profein partial cele	Homo sapiens mRNA for KIAA1337 protein partial cds	Homo saciens adaptor-related protein complex AD-4 angles cultural and angles AD-4 angles and angles AD-4 angles and angles AD-4 angles and angles AD-4 angles and angles AD-4 angles and angles and angles AD-4 angles and angles AD-4 angles and angles AD-4 angles and angles angles and angles and angles and angles and angles and angles and angles and angles and angles and angles and angles and angles angles and angles and angles and angles and angles and angles and angles and angles and angles and angles and angles and angles angles and angles and angles and angles and angles and angles and angles and angles and angles and angles and angles and angles angles and angles and angles angles and angles angles and angles angles angles and angles angles and angles and angles angles and angles angl	and 7001 xt NCI CGAP Part Home series and show WA CE and 247 of	H. sapiens ECE-1 gene (exon 17)	Homo sapiens HOXD13 gene for homeobox transcription factor, complete rule	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAMA) mRNA	Homo sapiens intersectin short isoform (ITSN) mRNA, complete cds	Homo sapiens T-oell tymphoma invasion and metastasts 1 (TIAM1), mRNA	Homo sapiens intersectin short isoform (ITSN) mRNA, complete ods	Human kappa-timmingolohidin saamilna nasantosaaa (Ob-22 4) saatata saamilna nasantosaa	Homo sepiens cysteine-rich reneat-centaining protein SS2 measures mBNA complete of	Homo saplens colladen, toe XII. alohe 1 (COI 1241) mRNA	Homo sapiens collagen, type XII, alpha 1 (COL12A1), mRNA	601497032F1 NIH MGC 70 Homo sapiens cDNA clone IMACE: 3898358 ff	601896173F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4125234 6	601896173F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4125234 5
Top Hit Detabase Source	5	LV.	Ę		İ	1 <del>2</del>	Į.	NT	Į.	T HUMAN		T HUMAN	Т	Т					LN	T HUMAN	Т										EST HUMAN	П	EST_HUMAN 6
Top Hit Acession No.	4557250 NT	1.0E-120 AB011399 1		7334	1.0E-120 AF056490.1					U134963.1	5032192	BF344378.1	F344378.1		9208.1	9208.1	3037758.1	1.0E-121 AB037758.1 N	1.0E-121 AF155156.2 N	Γ	-	-	11526176 NT	1.0E-122 AF114488.1 NT	11526176 NT	1.0E-122 AF114488.1 NT	M20707.1	167706.1	18424	11418424 NT	1.0E-122 BE906024.1 E	316170.1	316170.1
Most Similar (Top) Hit BLAST E Value	1.0E-120	1.0E-120	1.0E-120	1.0E-120	1.0E-120	1.0E-120	1.0E-120	1.0E-120	1.0E-121 Y	1.0E-121 A	1.0E-121	1.0E-121	1.0E-121 BI	1.0E-121	1.0E-121 Y1	1.0E-121 Y1	1.0E-121	1.0E-121	1.0E-121	1.0E-121	1.0E-121 X91937.1	1.0E-121	1.0E-122	1.0E-122	1.0E-122	1.0E-122	1.0E-122 M20707.1	1.0E-122 AF	1.0E-122	1.0E-122	1.0E-122 E	1.0E-122	1.0E-122 BF
Expression Signal	3.83	1.03	1.03	0.89	1.43	1.43	2.41	241	2.65	1.27	1.28	1.28	1.28	0.88	4.04	4.04	0.84	0.84	8.01	1.34	3.38	1.26	1.82	3.2	1.71	2.93	3.85	3.65	6.77	5.77	4.64	20.14	20.14
ORF SEQ ID NO:	11850	12143	12144	10368	14209	14210	14503	14504			10754	12584		12939			13494	13495	13607	14179	14781	14954				10931	11238	11722	11746	11747	11862	12511	12512
Exen SEQ ID NO:	6762	1031						9517	5150	5424	7728	7469	7469	7918	8024	8024	8467	8467	8600	9197	9801	9979	5324	5386	5406	5889	6201	6650	0299	6670	6770	7391	7391
Probe SEQ ID NO:	1770	2049	2049	3235	4232	4232	4527	4527	72	377	714	2501	2501	2899	3007	3007	3459	3459	3593	4204	4817	5008	265	334	320	84	1200	1854	1674	1674	1778	2420	2420

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Homo sepiens FYVE domein-containing dual specificity protein prospiratiese in victoria in total constructions of the containing dual specificity protein prospiratiese in victoria containing dual specificity protein prospiratiese in victoria containing dual specificity protein prospiratiese in victoria containing dual specificity protein prospiratiese in victoria containing dual specificity protein prospiratiese in victoria containing dual specificity protein prospiratiese in victoria containing dual specificity protein prospiratiese in victoria containing dual specificity protein prospiratiese in victoria containing dual specificity protein prospiratiese in victoria containing dual specificity protein	Homo sapiens amyloid beta (A4) precursor protein (protease nextr-II, Atzheimer disease) (APP), mRNA	ULAFERNO-BI-4-03-0-ULA NINI-MGC_30 India appeals Color MAGE:4153670 5	602018058F1 NCI CCAP Birdy from series CONA close MAGE 4153670 5	602018Uport NCL Conf. Billor Living against ASA(CD49	Homo septens diffunctional profession mithochandrial (mitofilla) (IMMT), mRNA	Homo sapiens little interiorate process, the little (PIPSK2B) mRNA, and translated	Homo sapiens phospinatodymostical reprosprints of the products products	Homo sapiens phosphetidylinositor 4-phosphate 3-ranase, type II, social II in companies products	SIND Flames (1997)	Homo saplens partiel mRNA for immunoglobulin kappa chain variable region (IGVN gene), sample Cavoz	Transactions of Annal Control of the Control of Cols	Human employee in (AMELY) sens 3 and of cds	Human garleng RABG-like protein (LOC51209), mRNA	Trails especially because the control of the contro	Home servers T-cell fumphome invasion and metastasis 1 (TIAM1) mRNA	Trough September 1 Cont. Surprise Investor and metastasts 1 (TIAM1) mRNA	Lorse services DNA for amidded precursor protein, complete cds	Truing seriens chromosome 21 segment HS21C046	THAT IN Stratagens schizo brain S11 Homo sapiens CDNA clone IMAGE:728719 5' similar to TR:G3004821	G300482 POL-REVERSE TRANSCRIPTASE HOMOLOG (RETROVIRAL ELEMENT);	Z81604.r1 Strategene schizo brain 311 Hours septicit Como Carlo III.  Gannas POL REVERSE TRANSCRIPTASE HOMOLOG (RETROVIRAL ELEMENT):	Human putative ribosomal protein S1 mRNA	Homo saplens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	Home seplens hypothetical protein (HSPC068), mRNA	Home saciens alucose transporter 3 gene, exons 9, 10, and complete cds	Homo serviens alucose transporter 3 gene, exons 9, 10, and complete cds	Homo sapiens mRNA for nucleolar RNA-helicase (not161 gene)	
Top Hit Datebase Source			L HUMAN	EST HUMAN	EST HUMAN																EST_HUMAN		FSI TOWNS			-	- 1	ž E	
	Ę	둗	EST	ES	ES	뒬	틸	Ž	N		Ę	킬	뉟		<u> </u>	킼		Z			<u> </u>			1 2		<u>z   2</u> 2	7	7	
· Top Hit Acesslon No.	\F264717.1	4502168 NT	1.0E-122 AW504645.1	1.0E-123 BF345274.1	3F345274.1	1.0E-123 AL163249.2	5803114 NT	4505818 NT	4505818 NT		1.0E-123 AJ388641.1	1.0E-123 M55419.1	M55419.1	55419.	1705962INI	1.0E-123 AL163280.2	4507500 NT	1	<u> </u>	AL163246.2	AA397551.1		_	AF-19303		_		_	140131712.1
Most Similar (Top) Hit BLAST E Value	1.0E-122 AF	1.0E-122	1.0E-122	1.0E-123	1.0E-123 BF	1.0E-123	1.0E-123	1.0E-123	4 00 422	1.05-123	1.0E-123	1.0E-123	1.0E-123 M	1.0E-123 M	1.0E-123	1.0E-123	1.0E-124	1.0E-124		1.0E-124	1.0E-124		1.0E-124	1.0E-124					1.0E-124
Expression Signal	0.98	2.44	1.49	2.05	2.05	5.43	8.48	4.18		4.18	2.66	1.75	1.75	1.75	2.82	0.93	2.41	2.41	1.29	2.33	5.13								4.44
ORF SEQ ID NO:	12810	14667		10807		11039	11046	l	l_	11261	11484	12134	12135			<u></u>	10334	5 10335	-	10530	40717			10798		1 10950	11373		11863
Exon SEQ ID NO:	787							١	1	6218	6425		L			L	L				2002	┸	5706	9 5771	9 5820	3 5911	_		12 6771
Probe SEQ ID NO:	2766	4600	4832	759	759	266	1006	1240	2	1219	1428	2042	2042	2042	2251	4326	266	36	272	482		289	682	749	8	893	1328	1328	1779

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	601491715F1 NIH_MGC_69 Hamo sepiens cDNA clane IMAGE:3833954 5	Homo saplens gene for B120, exon 11	Homo septens ATP-sensitive Inwardly rectifying K-channel subunit (KCNJ6/BIR1) gene, exon	Homo sapiens ATP-sensitive inwardly rectifying K-channel subunit (KCNJ6/BIR1) gene, exon	H.sapiens lactate dehydrogenase B gene exon 1 and 2 (EC 1.1.1.27) (and Johned CDS)	Homo sapiens T-cell lymphoma invasion and metastasis 1 (TIAM1) mRNA	Homo sapiens glutamata receptor, lonotropic, kainate 1 (GRIK1) mRNA	Homo sapiens gene for B120, excn 11	Human fibronactin gene extra type III repeat (EDII), extn x+1	Homo sapiens mRNA for KIAA1172 protein, partial cds	601577981F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3926685 5	Homo sapiens ALR-like protein mRNA, partfal cds	263207.s1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:486540 3' similar to gb:X65857_cds1 OLFACTORY RECEPTOR-LIKE PROTEIN HGMP07E (HUMAN);	Homo sapiens chromosome 21 segment HS21C010	Homo saplens KIAA0744 gene product; histone deacet/lase 7 (KIAA0744), mRNA	Homo sapiens Usurpin-alpha mRNA, complete cds	Homo sapiens Usurpin-alpha mRNA, complete cds	201009.rt Soares, fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone INAGE:428568 5	263307.s1 Soares_prognant_uterus_NbHPU Horino sapiens cDNA clone IMAGE:486540 3' strnllar to gb:X65857_cds1 OLFACTORY RECEPTOR-LIKE PROTEIN HGMP07E (HUMAN);	Homo sapiens inhibin, alpha (INHA) mRNA	Homo sapiens inhibin, alpha (INHA) mRNA	bb74f06.y1 NIH_MGC_12 Homo sepiens cDNA done MAGE:3046131 b' similar to III: Usbou4 Usbou4 ZINC FINGER PROTEIN.;	2453-07, s1 Soares_pregnant_uterus_NbHPU Homo septens CDNA clone INNAUE: 4865-40 3' sumilar to gb:X65657_cds1 OLFACTORY RECEPTOR-LIKE PROTEIN HOMPOTE (HUMAN);	Homo sapiens zinc finger protein ZNF287 (ZNF287), mRNA	Homo sapiens zinc linger protein ZNF287 (ZNF287), mRNA	601141152F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3140796 5	Homo sapiens CDC-like kinase (CLK) mRNA	Human laminin B1 chain gene, exon 20	H. sapiens gene for alphal-antichymotrypsin, exch 3	Homo sapiens hypothetical protein FLJ20048 (FLJ20046), mrvvn
Top Hit Database Source	EST_HUMAN	TN	NT	Z	N	7	7	NT	NT	TN	EST_HUMAN	N.	EST HUMAN	NT.	닐	N	N	EST HUMAN	EST_HUMAN	ĽN	TN.	EST_HUMAN	EST HUMAN	1	NT	EST_HUMAN	NT	IN	TN	NT
Top Hit Acession No.	378524.1	124069.1	684.1			4507500 NT	4504116 NT	024069.1	8178.1			Γ		163210.2	7662279 NT	1.0E-125 AF015450.1				4504696	4504696 NT	1.0E-125 BE018009.1	042813.1	11425114	11425114 NT	1.0E-125 BE315412.1	4758007 NT	M61936.1	X68735.1	8923056 NT
Most Similar (Top) Hit BLAST E Value	1.0E-124 BE	1.0E-124 AB(	1.0E-124 S78	1.0E-124 S78684.1	1.0E-124 X13794.1	1.0E-124	1.0E-124	1.0E-124 AB	1.0E-124 M1	1.0E-125 AB032998.1	1.0E-125 B	1.0E-125 AF264750.1	1 OF-125 AA042813 1	1.0E-125 AL	1.0E-125	1.0E-125	1.0E-125 /	1 0F-125 /	1.0E-125	1.0E-125	1.0E-125	1.0E-125	1 0E-125 AA	1.0E-125	1.0E-125	1.0E-125	1.0E-128	L		Ш
Expression Signal	1.32	1.32	0.74	0.74	1.19	0.88	0.81		1.39	11.96	4.13	1.95										0.98								
ORF SEQ ID NO:	12096					L			L		10065						L					l			L	1		1		
Exon SEQ ID NO:	6992	L			8574		1		L		1_	L				ı	1_	L	1				L	8//1				L		
Probe SEQ ID NO:	2009	2388	3410	3410	3564	3804	3962	4601	4778	317	423	718	3	883	44.28	1775	1775		2431	2520	2520	200		3/68	4410	4484	785	768	2 8	2283

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Single Exon Probes Expressed in HBL100 Cells

SEON D. 12.         CRG SEON D. PHOL.         Robert Standard (Top)-IRI Top-IRI No. Bushase         Top-IRI Descriptor		_	_	_	-	_	-	_	_	_	_	-	-	_	_			_	_	<u>N</u>		Ļ	4	<u> </u>	7	40	_1_	**	Ц.,	1 I)	1 1:	1
Exon DORF SEQ Boynession NO: Signal DIONO:         Most Similar Signal DIONO:         Most Similar Signal DIONO:         Most Similar Signal DIONO:         Top Hit Acession DIONO: <td>Top Hit Descriptor</td> <td>lamo sapiens hypodhetical protein FL (20048 (FL J.) m.R.N.A.</td> <td>Iomo saciens RAN binding protein 2 (RANRP2) mRNA</td> <td>D72003.1 Stratagene pencress (#037208) Home seniors CDNA close MAACE SO2420 E</td> <td>D72c03.r1 Strategene parkreas (#937208) Homo saviens cDNA close MAGCE-rangon kr</td> <td>sapiens DNA for liver cytochrome b5 bsaudocene</td> <td>Idmo sapiens death receptor 6 (DR6), mRNA</td> <td>Iomo sapiens collagen type XI alpha-1 (COL11A1) cene exm 63</td> <td>Iomo saplens collagen the XI alpha-1 (COL 1141) cane com 63</td> <td>x78c06.r1 Soares melanocyte ZNbHM Homo saciens cDNA clone IMAGE: 267850 6</td> <td>lano sapiens mRNA for casein kinase l'ensilon, camplete cas</td> <td>Iomo sapiens mRNA for casein kinase i epsilon, complete cds</td> <td>Iomo sapiens mRNA for casein kinase l'epsilon, complete cds</td> <td>lomo sapiens mRNA for casein kinase I epsilon, complete cds</td> <td>omo sapiens DNA for amyloid precursor protein, complete cds</td> <td>omo sapiens DNA for amyloid precursor protein, complete cds</td> <td>omo sapiens intersectin short Isoform (ITSN) mRNA, complete ods</td> <td>omo sapiens lost on trensformation LOT1 mRNA, complete cds</td> <td>omo sapiens ubiquitin specific protease 8 (USP8) mRNA</td> <td>omo sapiens leukocyte Immunoglobulin-like receptor, subfamily A (with TM domain), member 1 (LILRA1), IRNA</td> <td>iomo sapiens feukocyte immunoglobulin-like receptor, subfamily A (with TM domain), member 1 (LILRA1), PNA</td> <td>omo sapiens ribosamel protein L26 (RPI 26) mRNA</td> <td>omo sapiens adlican mRNA, complete ods</td> <td>uman mRNA for cytokeratin 18</td> <td>omo sapiens intersectin short isoform (ITSN) mRNA, complete cds</td> <td>U80908 y1 Schneider fetal brain 00004 Homo sapiens CDNA done IMAGE:2782594 &amp; similar to R.C.15170 Q15170 TRANSCRIPTION FACTOR S.II.RELATED PROTEIN complex element MFR22</td> <td>petitive element;</td> <td>отто sapiens neuroblastoma-amplified protein (LOC\$1594), mRNA</td> <td>omo sapiens neuroblastome-emplified protein (LOC51594), mRNA</td> <td>omo sapiens RAD1 (S. pombe) homolog (RAD1) mRNA, and translated products</td> <td>omo sapiens chromosome 21 segment HS21C068</td> <td>omo sapiens Ring1 and YY1 binding protein (RYBP), mRNA</td>	Top Hit Descriptor	lamo sapiens hypodhetical protein FL (20048 (FL J.) m.R.N.A.	Iomo saciens RAN binding protein 2 (RANRP2) mRNA	D72003.1 Stratagene pencress (#037208) Home seniors CDNA close MAACE SO2420 E	D72c03.r1 Strategene parkreas (#937208) Homo saviens cDNA close MAGCE-rangon kr	sapiens DNA for liver cytochrome b5 bsaudocene	Idmo sapiens death receptor 6 (DR6), mRNA	Iomo sapiens collagen type XI alpha-1 (COL11A1) cene exm 63	Iomo saplens collagen the XI alpha-1 (COL 1141) cane com 63	x78c06.r1 Soares melanocyte ZNbHM Homo saciens cDNA clone IMAGE: 267850 6	lano sapiens mRNA for casein kinase l'ensilon, camplete cas	Iomo sapiens mRNA for casein kinase i epsilon, complete cds	Iomo sapiens mRNA for casein kinase l'epsilon, complete cds	lomo sapiens mRNA for casein kinase I epsilon, complete cds	omo sapiens DNA for amyloid precursor protein, complete cds	omo sapiens DNA for amyloid precursor protein, complete cds	omo sapiens intersectin short Isoform (ITSN) mRNA, complete ods	omo sapiens lost on trensformation LOT1 mRNA, complete cds	omo sapiens ubiquitin specific protease 8 (USP8) mRNA	omo sapiens leukocyte Immunoglobulin-like receptor, subfamily A (with TM domain), member 1 (LILRA1), IRNA	iomo sapiens feukocyte immunoglobulin-like receptor, subfamily A (with TM domain), member 1 (LILRA1), PNA	omo sapiens ribosamel protein L26 (RPI 26) mRNA	omo sapiens adlican mRNA, complete ods	uman mRNA for cytokeratin 18	omo sapiens intersectin short isoform (ITSN) mRNA, complete cds	U80908 y1 Schneider fetal brain 00004 Homo sapiens CDNA done IMAGE:2782594 & similar to R.C.15170 Q15170 TRANSCRIPTION FACTOR S.II.RELATED PROTEIN complex element MFR22	petitive element;	отто sapiens neuroblastoma-amplified protein (LOC\$1594), mRNA	omo sapiens neuroblastome-emplified protein (LOC51594), mRNA	omo sapiens RAD1 (S. pombe) homolog (RAD1) mRNA, and translated products	omo sapiens chromosome 21 segment HS21C068	omo sapiens Ring1 and YY1 binding protein (RYBP), mRNA
Exon NO:         ORF SEQ ID NO:         Expression Signal         Most Similar (Lop) Htt         Top Htt Acession Plane           7256         12378         1.21         1.0E-126         8923056           7488         12607         2.83         1.0E-126         8923056           7488         12607         2.83         1.0E-126         8382079           8017         13030         64.41         1.0E-126         AA160706.1           8072         13559         0.75         1.0E-126         AA160706.1           8578         13559         0.75         1.0E-126         AA160706.1           8578         13559         0.75         1.0E-127         AA100706.1           8627         14619         0.88         1.0E-127         AA100706.1           85236         10247         8.8         1.0E-127         AA101108.1           85236         10248         8.8         1.0E-127         AA10148.1           65236         10248         8.8         1.0E-127         AA10148.1           6536         10248         8.4         1.0E-127         AB024597.1           6536         10248         8.4         1.0E-127         AB024597.1           6530         1	Top Hit Defabase Source			T HUMAN	Г					Г																						
Exon NO:         ORF SEQ Expression ID NO:         Expression Signed Sign	Top Hit Acession No.	8923056	6382078	Γ			7657038				Г						Γ		4827053	5803065	FROTORE	4506620	Π					7706239	7706239 N	4508384	NL163268.2	6912639 N
Exam NO:         ORF SEQ ID NO:         Express Signs           7259         12378         Signs           7488         12607         6           8017         13030         6           8077         13659         6           8077         13659         6           8578         13584         6           8677         14619         6           8678         10247         6           5236         10247         6           5236         10248         6           5236         10247         6           5236         10248         6           5236         10247         6           5236         10248         6           6648         10248         6           6648         1720         1           7117         12231         3           7256         12620         1           8613         13621         1           8613         13621         1           8462         14441         2           8488         14513         2	Most Similar (Top) Hit BLAST E Value	1.0E-126	1.0E-126	1.0E-126	1.0E-126			1.0E-126	1.0E-126	1.0E-126	1.0E-127	1.0E-127	1.0E-127	1.0E-127	1 0F-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127		1.0E-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127	1.0E-127						
Exan SEQ ID ID NO: 17269 17488 8017 8017 8017 8017 8017 8017 8027 8027 8027 8027 8028 6038 6038 6038 6038 6038 6038 6038 603	Expression Signal	1.21	2.83	64.41	54.41	0.75	2.04	0.98	0.98	1.53	8.83	8.83	8.4	8.4	1.45	1.45	1.36	1.51	1.92	2.09	2.09	38.06	2.58	16.49	0.82		17.44	22.49	22.49	5.1	3.63	1.21
"" "	ORF SEQ ID NO:	12378	12607	13030	13031	13559	13584	14619	14620	14654	10247	10248	10247	10248	10342	10343	10930	10959	11720	12099	12100	12231	12374	12620	13621		13725	14110	14111	14441		14513
Probe SEQ ID NO: 2283 2299 2299 3544 3574 4642 4642 4687 177 177 177 177 177 177 177 177 177 1		7259	7488	8017		·		9627	9627	9672	5236	5236	5236	5236	2330	5330	5888	5922	6648	9889	6996	7117	7256	7500	8613		8725	9127	9127	9462	9488	8527
	Probe SEQ ID NO:	2283	2521	2999	2999	3545	3571	4642	4642	4687	13	Ē	172	172	271	27.1	870	902	1652	2013	2013	2137	2280	2535	3606		3721	4132	4132	4472	4498	4537

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Single Exon Probes Expressed in HBL100 Cells

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Single Explicationes Expressed in ABL100 Cells	Top Hit Descriptor	601278127F1 NIH MGC 20 Homo sapiens cDNA close IMACE-3848822 #	Human FAU1P oseudocene, trinucleotide renderf reclines	Human FAU1P pseudopene trining inches resistant resistant	Homo saplens ribosomal protein S2 (RPS2) mRNA	Homo sablens chromatin enacific framendation absention forthe 440 D.	Homo sablens mRNA for KIAA1247 protein, narkel ods	Homo saplen's prospero-related homeobox 1 (PROX1) mRNA	insulin-like growth factor bridging protein-2 flumen placents. Genomic 1019 at exempts 2 4 41	instilin-like growth factor binding brothing. Thuman placents. Generally, 1010 at animars 4 of 1	Novel human mRNA containing Zinc finger C2H2 type domains	Homo sapiens glutathione S-transferase theta 2 (GST12) and glutathione S-transferase theta 1 (GSTT1) genes, complete cds	Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1) denses, complete eds	Homo sabiens zinc finder profein 76 (Appressed in feetle) (7NE76)	ZINC FINGER PROTEIN HZF10	ZING FINGER PROTEIN HZF10	ZINC FINGER PROTEIN HZF10	Homo saplens A kinase (PRKA) anchor protein 9 (AKAP9), mRNA	Homo sapiens A kinase (PRKA) anchor protein 9 (AKAP9), mRNA	Homo sapiens mRNA for KIAA1459 protein, partial cds	CMYA5 Human cardiac muscle expression library Homo saplens cDNA clone 4151935 similar to CMYA5 Cardiomyopathy essociated gene 5	CMYAS Human cardiac muscle expression library Homo sepiens cDNA clone 4151935 similar to CMYA5	Carquentyopatry associated gene 5	From Septembrily Institute of court (Tiest C.244), mitaly 601121895F1 NIH MGC 20 Home serviews of MA above MAACE 39 Agree F1	601121995F1 NIH MGC 20 Hamo sapiens CDNA clone MAGE:3348388 61	Human gene for catalase (EC 1.11.1.6) exon 9 mapping to chromosome 11, band p13	Homo sapiens RET finger protein-like 1 antisense transcript, partial	601343016F1 NIH_MGC_53 Homo septens cDNA clone IMAGE:3685466 5	601343016F1 NIH_MGC_63 Hamo sapiens cDNA clone IMAGE:3685466 5'	Homo sapiens retinol dehydrogenase homolog isoform-1 (RDH) mRNA, complete cds
Second Lines	Top Hit Database Source	EST HUMAN	Į.	Z	L		L'X	F	TN TN	N	N L	_ <del>F</del>	- <u>F</u>	1	SWISSPROT	SWISSPROT	SWISSPROT	77	<u></u>	Į,	EST HUMAN		ES HOMAN	EST HIMAN	EST HUMAN	Į.	N	EST_HUMAN	EST_HUMAN	N
BISHIS	Top Hit Acession No.	E385617.1	102523.1		4506718	11437455INT	1.0E-128 AB033073.1	26673			1.0E-129 AL096880.1	1.0E-129 AF240786.1	1.0E-129 AF240786.1	8522				5032230	5032230 NT	1.0E-129 AB040892.1	N755254.1		1.0E-128 AW (55254.1 ES	1.0E-130 BE276192 1	275192.1		1.0E-130 AJ010230.1			-240698.1
	Most Similar (Top) Hit BLAST E Value	1.0E-128 B	1.0E-128 U	1.0E-128 U02523.1	1.0E-128	1.0E-128	1.0E-128	1.0E-128	1.0E-129 S37722.1	1.0E-129 S37722.1	1.0E-129 /	1.0E-129	1.0E-129	1.0E-129	1.0E-129 Q14585	1.0E-129 Q14585	1.0E-129 Q14585	1.0E-129	1.0E-129	1.0E-129 /	1.0E-129 A)	4 00 400	1 0E-128	1.0E-130 F	1.0E-130 BI	1.0E-130 X04092.1	1.0E-130	1.0E-130 E	1.0E-130 BE564219.1	1.0E-130 Ai
	Expression Signal	4.57	12.06	12.06	127.93	4.72	1.28	4.83	1.18	1.25	3.33	1.56	1.58	5.19	1.71	1.71	1.71	1.03	1.03	1.96	2.16	9,0	40.4	31.59	31.59	2.05	5.31	1.17	1.17	0.78
	ORF SEQ ID NO:	10506	12104		12243		13344	14506	10469	10469	11750	11755	11756	11890	13082	13083	13084		13995	14026	14135	44438	10163	11693	11694			12844	12845	13520
	Exam SEQ ID NO:	5494		7001	7126	7349			5447	5447	9299	6681	6681	6229	8072	8072	8072	9006	9006	9036	9153	0153	5153	6624	8624	6923	7655	7829	7829	8505
	Probe SEQ ID NO:	457	2018	2018	2147	2377	3310	4529	122	410	1680	1685	1685	1809	3022	3055	3055	4010	4010	4040	4158	4459	32	1627	1627	1837	2698	2809	5808 7808	3497

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	MASACATES NET MODE SO Hammer and the second	SA12/2016 A MILL 1400 TO THE SEPTENS CLINA CICHO IMAGE: 3685466 6	COLOMBINITATION OF THE WILL AND SEPTEMBERS CLAVE CHONE INVICE: 3665466 5	Himm T. All records 07 4-15 22 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	CMA CNOVAS, 180200 511 002 CNINGS 11	RCI-CIO318-201400-031-102 CANDU43 India September CLINA	RO-CT0318-201199-031-011 CT0318 Home septems CDNA	## ## ## ## ## ## ## ## ## ## ## ## ##	278604.17 Soars NH-IMPLY ST HOME septents CDNA done IMAGE:667590 5' similar to TR:G222811	Homo saplems checkmaint en manages 4 (Clares)	Homo seciens DCRR mRNA martiel An	Homo sepiens DCRR1 mBNA marial Ac	Homo sapiens beta-tubulin mRNA complete Ac	Homo saplens Cole42 effection motein 2 (CED2) - Data	Human heparin cofactor II (HCF2) gene grove 1 through 6	Homo sepiens RNA-binding protein S1, serine-rich domain (RNDS4), mRNA	Homo sapiens mRNA for multidatio resistence number 3.74 (2007)	Homo sapiens mRNA for multidua resistence protein 3 (ARCC3)	HUM516H08B Human placenta polyA+ (TFuiwara) Homo sapiens chwa claws CEN Estatos et	HUM516H08B Human placenta polyA+ (Truiwara) Homo sapiens con Achae CEN Exemple 5	Human ribosomal protein L7 (RPL7) mRNA, complete cds	cr48e07.x1 Jia bone marrow stroma Homo sapiens cDNA clone HBMSC cr48e07.31	cr48e07.x1 Jia bone marrow stroma Homo saplens cDNA clone HBMSC cr48e07.31	Homo sapiens protein tyrosine phosphatase, non-receptor type substrate 1 (PTPNS1) mRNA	Homo sepiens protein tyrosine phosphatase, non-receptor trae substrate 1 (PTPNS1) mBNA	Homo sapiens protein tyrosine phosphatase, non-receptor tyros substrate 1 (PTPNS4) - INNA	Homo saplens protein tyrosine phosphalase, non-receptor type substrete 1 (PTPNS-1) mRNA	Homo sephens amilioride binding protein 1 (amine oxidase (copper-containing)) (ABP1), nuclear gene	Make in the control of the control o	Homo serviens actin here (ACTR) mRNA	mis replyoned 4 banded (1014) DAIA	THE PAY TO THE TOTAL OF THE TANK OF THE TA
Top Hit Database Source	EST HIMAN	EST HIMAN	EST HIMAN	L L	HIMAN	Т	Т		Т						,			L L	HUMAN	EST_HUMAN	П		EST HUMAN		IN	L L	- LN					
Top Hit Acessian No.	E564219.1	E564219 4		T	2	Τ	1.0E-130 AW363299.1			35136	083327.1		=	7882		3857825	7151.2						0.0E+00 AW069534.1 E	4758977 N	4758977 N	4758977	4758977	4501850 NI	AFOAAAA NIT	5016088 NT	R9277 1	
Most Similar (Top) Hit BLAST E Value	1.0E-130 B	1.0E-130	1.0E-130	1.0E-130	1.0E-130	1.0E-130	1.0E-130	0.0E+00	0.0E+00	0.0E+00	0.0E+00 D83327.1	0.0E+00 D83327.1	0.0E+00	0.0E+00	0.0E+00 M58600.1	0.0E+00	0.0E+00 Y1	0.0E+00 Y1	0.0E+00 D78804.1	0.0E+00 D78804.1	0.0E+00 L16558.1	0.0E+00 /	0.0E+00 /	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.05+00.0	00+100	0.0E+00	0.0E+00/LIRG277 4	7
Expression Signal	5.51	5.51	1.54	1.25	9.21	1.3	1.3	1.8	1.8	1.14	5.04	5.04	107.34	1.77	0.74	24.17	4.3	4.3	2.89	2.89	54.73	11.31	11.31	2.95	2.95	1.73	1.73	7.0	50.8	77.93	45.78	
ORF SEQ ID NO:	12844	12845	13841	13947	14383	14880	14881	10069	10070	10074	10085	10086	10091	10101	10104	10109	10136	10137	10141	10142	10143	10146	10147	10161	10162	10161	10162	10168		10177	10180	
Exen SEQ ID NO:	7829	7829	8835	8958	8338	9903	9903	5085	5085	5088	5101	5101	5107	5115	5117	5121	5138	5138	5140	5140	5141	5143	5145	5152	5152	5152	5152	6157	5158	5167	5170	
Probe SEQ ID NO:	3681	3681	3833	3960	4408	4926	4926	4	4	8	2	72	72	33	37	4	8	8	8	8	8	8	3 1	₹	7	F	=	8	٣	8	8	

Page 173 of 209 Table 4 Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	HA1347 Human fetal liver cDNA library Homo sapiens cDNA	Homo sepiens mRNA for KIAA1363 protein, pertial cds	1838-05-17 NCI_CGAP_UM Homo sepiens d'ONA cione IMAGE:2230833 3° similer to TR:099551 099561 MITOCHONDRIAL TRANSCRIPTION TERMINATION FACTOR PRECURSOR.;	1838-05.rd NCI_CGAP_UM Homo sepiens CDNA clone INAGE:2230833 3' similer to TR:Q98551 Q99551 MITOCHONDRIAL TRANSCRIPTION TERMINATION FACTOR PRECURSOR.;	yy01h09,r1 Scares melanocyte 2NbHM Homo sepiens cDNA clone IMAGE:270017 51	yy01h09.r1 Soares melanocyte ZNbHM Home septens cDNA clone IMAGE:270017 5	Homo sapiens neuropilin 2 (NRP2) mRNA	Homo sapiens polymerase (RNA) II (DNA directed) polypeptide A (220kD) (POLR2A) mRNA	Homo sapiens polymerase (RNA) II (DNA directed) polypeptide A (220kD) (POLR2A) mRNA	ya83g04.r2 Stratagene fetal spleen (#937205) Homo sapiens cDNA clone IMAGE:68310 6	ya83g04.r2 Stratagene fetal spleen (#937205) Homo saplens cDNA clone IMAGE:68310 5	Homo sapiens heterogeneous nuclear ribonucleoprotein A1 (HNRPA1) mRNA	601460375F1 NIH_MGC_66 Hamo sapiens cDNA clane IMAGE:3863803 6	Homo sapiens heterogeneous nuclear ribonucleoprotein A1 (HNRPA1) mRNA	Homo sapiens serine palmitory transferase, subunit II gene, complete cds; and unknown genes	601174270F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3529864 5	601174270F1 NIH_MGC_17 Hamo septens cDNA clane IMAGE:3529864 5'	zd62b05.r1 Soares_fetal_heart_NbHH19W Homo sepiens cDNA clone IMAGE:345201 6" similar to	gb:X16282_cds1 ZINC FINGER PROTEIN CLONE 647 (HUMAN);	QV3-HT0457-140200-088-d04 HT0457 Homo sapiens cDNA	QV3-HT0457-140200-088-d04 HT0457 Homo sapiens cDNA	Homo sapiens zinc finger protein mRNA, complete cds	Homo sapiens chromosome 21 segment HS21C002	Homo sapiens chromosome 21 segment HSZ1 W02	bb24e(2.y1 NIH_MGC_14 Homo sapiens cDNA clone IMAGE:2963854 5 straitar to WP:75/A10A.Z	CEZZOST ;	bb24e12.y1 NH_MGC_14 Homo septens culva done invave=.25005094 o suimm to viri : 10101504. CE22631;	Homo sapiens mRNA for KIAA0784 protein, partial cds	Homo sapiens mRNA for KIAA0784 protein, partial ods	Homo sapiens mRNA for KIAA0784 protein, partial cds	Homo sapiens mRNA for KIAA0784 protein, partial cds	
Top Hit Database Source	EST_HUMAN	IN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	41	۲۲	EST_HUMAN	EST_HUMAN	77	EST HUMAN	L/L	NT	EST_HUMAN	EST_HUMAN		EST HUMAN	EST_HUMAN	EST_HUMAN	NT	L	NT		EST HUMAN	EST_HUMAN	N	N F	K	¥	
Top Hit Acession No.	4743.1	0.0E+00 AB037784.1					4505458	4505938 NT	4505938 NT			4504444	0.0E+00 BF036881.1	4504444 NT			0.0E+00 BE295973.1		3973.1	162832.1	62832.1	244088.1	63202.2	63202.2		0.0E+00(BE018970.1	BE018970.1	018327.1	318327.1	Γ	0.0F+00 AB018327.1	
Most Similar (Top) Hit BLAST E Value	0.0E+00 AI11	0.0E+00	0.0E+00	0.0E+00	0.0E+00 N36040.1	0.0E+00 N36040.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00 T56945.1	0.0E+00 T56945.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00 AF1	0.0E+00	0.0E+00		0.0E+00 W7	0.0E+00 BE	0.0E+00 BE	0.0E+00		0.0E+00 AL1		0.0E+00	0.0E+00		1_	1_		١.
Expression Signal	2.36	1.89		1.65	3.4	3.4	0.7	4.8	4.8	1.18	1.18	7.07	3.22	143.39	0.82	6.28	1.45		6.65	2.09	2.09	3.77	15.16	15.16		9.67	79 6	423				
ORF SEQ ID NO:	10187	10188	10197	10197	10198	10199	10204	10212	10213	10220	10221		10234		10237	10238			10239						L	10253	10254					
Exon SEQ ID NO:	5177	5178	5188	5188	7690	7690	5191	5197	5197	5205	5205	5218	5222	5224	5227	5229	5229		5230		L	L	5235			5243	<u> </u>		$\perp$		20,02	
Probe SEQ ID NO:	8	5	116	117	178	138	121	131	134	\$	140	152	156	158	19	58	164		165	166	166	167	170	170		180	1	30,	207	2 3	00	180

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Table 4

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Single Exon Probes Expressed in HBL100 Cells	Top Hit Descriptor		Human gamma-cytoplasmic actin (ACTGP9) pseudogene	From 6 septems CTCL fumor entigen self43 mRNA, complete cds	nome sapiens CICL fumor entigen self-3 mRNA, complete cds	From septens chromosome X MSL3-2 protein mRNA, complete cds	rene septeta culturiosanie X MSL3-2 protein mRNA, complete ods tq04f08.47 NCI_CGAP_Ut3 Homo septens cDNA done IMAGE:2207847.3' similar to gb:.103191 PROFILIN I I/HI MANY	tq04f08x1 NCL_CGAP_UB Homo sepiens cDNA done IMAGE:2207847 3' similer to ob:.103181 PROFII IN 1	יייין ייין יייין י	Inomo saprens DNA mismatch repair protein (MLH3) gene, complete cds	Tours squers mosame protein L31 (RPL31) mRNA	Induo sapients I AUA1 protein mRNA, complete cds	nomo septens mKNA for KIAA0721 protein, pertiel cols	Hamo sapiens mRNA for KIAA0721 protein, partial cds	Mus musculus testis-specific protein, Y-encoded-like (Tspyl), mRNA	nomo sapiens NS1-associated profein 1 (NSAP1) mRNA	Indiano sapiens chromosome 21 segment HS21C001	rionio saprens ciriomosome 21 unknown mRNA	n.sapiens mixivA for interferon alpha/beta receptor (long form)	Home septens difference 21 unknown mRNA	Homo serviens 1 cell lymphome invesion and metastasis 1 (TIAM1) mRNA	Homo service 1 Con minoroma my secon and metastasis 1 (TIAM1) mRNA	Homo septeme 17/0004 Data	Homo sapiene DODD4	Home capiese DOOD - DAILY	11 2 CT0031-184100 App Bras CTANAS 11			Homo serving mRNA 62 KIA A 1010		Homo services interval of NAMINIE protein, partial cds	nsferase, phosphoribosylgiycinamide synthetase, mRNA
Exon Probes	Top Hit Database Source	- 14	E L	ž į	1	LZ	EST HUMAN	POT LICENSAN	NAMOR IN	F		IN LIVE	TIV	12	1	LN LN						 	L	L	5	EST HUMAN	-1	-	LN	L	TN	<u>_</u> _
L	Top Hit Acession No.	DSORFO 4					AI587308.1	AI587308 4	AF195658 1	4506632 NT	AF13200	T	AB018264 1	6678444		Al 183201 2	T	T	AF231919 1	4507500	4507500 NT	7706028 NT	D83327.1	383327.1	383327.1	W845293.1	7029	4557029 NT	0.0E+00 AB028942.1	Γ	6728	4503914 NT
Mant Cimils	(Top) Hit BLAST E Value	0.0E+00	0 0F+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00/	0.0E+00/	0.0E+00	0.0E+00
	Expression Signal	261.78	3.04	3,01	28.3	28.3	57.42	57.42	1.9	99.79	15.17	3.41	3.35	4.92	16.43	17.29	3.82	1.5	6.35	1.48	1.48	2.14	2.81	2.43	2.43	0.88	5.4	5.4	5.25	4.98	48.48	2.02
	ORF SEQ ID NO:	10271	L		10279	П	10286	10287	10289			10293	10293	10294	10309	-	10315	10318		10336	10337	10339		10352	10353		10360	10361	10371	10372		10373
	SEQ ID NO:					5265	7715	7715	5275	5277	5278	5283	5283	5284	5298	2300	5305	5307	5315	5326	5326	5328	2338	5339	5339	5340	5348	5348	5359	5360	7718	5361
	SEQ ID NO:	194	199	199	201	201	210	210	212	24	215	22	221	222	236	238	245	247	255	267	287	88	8	281	281	282	291	781	302	303	304	305

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Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	2/18008.1 Source NIHMPu S1 Home senions of NA claw MACE-75/2004 F	Homo saplens SON DNA binding protein (SON) mRNA	Homo saplens SON DNA binding protein (SON) mBNA	Homo sablens intersectin short is from (TSN) mith a complex con-	TRANSCRIPTION REGULATOR PROTEIN BACH (BTR AKIN CNC HOLLY) OC 17 (UASSAN)	TRANSCRIPTION REGULATOR PROTEIN BACH! (BTB AND CNC HOMOLOG 1) (HAZSOS)	Homo sapiens hormonally uprequiated neu tumor-associated triass (HI MK)	Homo sapiens hormonally uprequiated neu tumor-associated kinasa (HI NiK) mDNA	Homo sapiens myeloid/lymphold or mixed-lineage feukemia (trithorax (Drosophila) homolog); translocated to, 4 (MLLT4) mRNA	Homo sapiens moesin (MSN), mRNA	omo saplens X-bra hindina ny fahan 4 (XBD4) mBNA	Human zinc finger brothen zin31 (#34) mRNA nortical colo	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens chromosome 21. unknown mRNA	Homo sapiens chromosome 21 unknown mRNA	Homo sapiens Tidel Ivanihama investim and materiacie 4 (TIANA) and NA	Homo sapiens GA-binding ordein transcription factor, ainthe submit (2010) (CABBA)DAIA	Human mRNA for KIAA0184 oene, partial cols	Human mRNA for KIAA0184 owns partial cds	Homo sapiens T-cell Imphoma Invasion and metastasis 1 (TIAM1) mRNA	AU134963 PLACE1 Homo saplens cDNA done PLACE1000899 51	Homo sapiens mRNA for KIAA1019 protein, partial cds	qy81h05x1 NCI_CGAP_Bm25 Homo septens cDNA clone MAGE:2018457 3' similar to gib:X64199 PHOSPHORIBOSYLAMINEGLYCINE LICASE / HINAAN:	RC2-CT0320-300100-016-409 CT0320 Homo seniens cDNA	Homo saplens IgG Fc binding protein (FC/GAMMA IBP) mRNA	Homo sapiens (gG Fc binding protein (FC(GAMMA)BP) mRNA	Homo sapiens IgG Fc binding protein (FC(GAMMA)BP) mRNA	Homo sapiens IgG Fc binding protein (FC(GAMMA)BP) mRNA	Homo saplens IgG Fc binding protein (FC(GAMMA)BP) mRNA	Homo sepiens IgG Fc binding protein (FC(GAMMA)BP) mRNA	Homo sepiens IgG Fc binding protein (FC(GAMMA)BP) mRNA	H. sapiens gene for RNA pol II largest subunit, exons 23-29
	Top Hit Database Source	EST HUMAN	Þ	¥	Į.	SWISSPROT	SWISSPROT	N.	Z E	 	17		LN N	Į.	LN L	N.	5	Þ	ΤZ	F	F	EST_HUMAN	77	EST HUMAN	EST HUMAN	7	5	F-7	<b></b>	7	7	47	L
B	Top Hit Acession No.	AA480002.1	4507152 NT	4507152INT	AF114488.1		014867	7657213 NT	7657213 NT	5174574 NT	4505256 NT	4827057 NT	U71600.1	AF231919.1	4F231919.1	F231919.1	4507500	4503854 NT	J80006.1		507500		0.0E+00 AB028942.1	0.0E+00 Al363014.1	0.0E+00 AW754180.1	4503680 NT	4503680 NT	4503680 NT	4503680 NT	4503680 NT	4503680 NT	4503680 NT	74870.1
	Most Similar (Top) Hit BLAST E Vatue	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0,0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
	Expression Signal	2:32	20.77	22.16	. 22	2.91	2.91	3.75	2.04	2.33	1.6	10.77	2.29	2.65	2.65	3.33	1.13	1.36	2.25	1.75	1.21	2.21	7.9	7.2	3.05	1.12	2.02	2.02	1.2	1.19	1.19	3.59	3.82
	ORF SEQ ID NO:						10388	10389	10389	10401	10402	10406	10411	10415	10416	10417	10419	10423	10424	10424	10426	10437	10479	10480	10444	10447	10448	10449	10450	10451	10452	10453	10454
	ш W ~	5362	සිසි	5363	5367	5379	5379	5380	9380	5394	5395	5398	5401	5405	5405	7719	5407	5410	5411	5411	5413	5422	5461	5462	5429	5431	5432	5432	5433	5434	5434	5435	5436
	Probe SEQ ID NO:	308	307	308	312	325	325	326	327	342	343	346	349	354	354	355	357	360	361	362	364	375	386	387	391	394	395	395	396	397	397	398	399

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	H serviens game for BNA mail I learned in the serviens game for BNA mail in the services and services are services and services and services and services are services and services and services are services and services and services are services and services and services are services and services and services are services and services and services are services and services are services and services and services are services and services are services and services are services and services are services and services are services and services are services and services are services and services are services and ser	H saniens gave for BNA and II have a second in the second	Hearing ages for DNA - 1117 - 1177 -	Home seniors therefore the contract of the con	Value seprets incoming protein ANIB Home seprets in 1995 Ania Ania Ania Ania Ania Ania Ania Ania	Homo sapiens phosphoribosylgiyahamida formytransferase, phosphoribosylgiyahamida synthetase,	Home carions the contraction of the Contraction of	Lymp and a microwing protein of (1700) mkink	Homo series CON DNA Nichama Series (CON)	Homo saplens SON DNA bioding protein (SON) — DNA	Mus musculus tracerted SON sector (Son)	Homo sapiens chromosome 21 segment 1624 CAN	Homo sablens interferm general 10210001	EST27054 Cerebellum II Homo sociene e Divis St	601111520F1 NIH MGC 16 Home series and a language for the language of the lang	Homo saplens Shudooutnatamina (sendrala) 48 / 140/20	Homo sapiens 5-hydroxydrynamine (serytonin) receptor 18 (117018) mrthy	Homo saplens keratin 18 (KRT18) mRNA	Homo sapiens keratin 18 (KRT18) mRNA	Homo sapiens chromosome 21 segment HS21Ch48	Homo sapiens chromosome 21 segment HS21C046	Homo saplens chromosome 21 segment HS21C046	Homo sapiens mRNA for KiAA1209 protein, partial cds	AU132898 NT2RP4 Homo sapiens cDNA clone NT2RP4000837 5	601274951F1 NIH MGC 20 Homo saplens cDNA clone MAGE 3815758 5	PM0-DT0065-130400-002-c06 DT0065 Hamp sacients aDNA	Novel human gare mapping to chomosome 1	Homo sapiens PC328 protein (PC326), mRNA	IL2-FT0159-070800-120-F07 FT0159 Homo sapiens oDNA	Homo sapiens chromosome 21 segment HS21C010	QV2-BT0635-160400-142-h05 BT0635 Homo sepiens cDNA	601764858F1 NIH_MGC_53 Hamp saplens cDNA clane IMAGE:3998998 5	Homo sepiens mRNA for KIAA1476 protein, partial cds
	Top Hit Database Source	LN	Į.	Z	5	EST HUMAN		1	Į	Ę	Į.	Z	Z	Į.	EST HUMAN	EST HUMAN	Z	Į,	12	77	N-	N-	LN PA	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	N	5	EST HUMAN	IN	EST_HUMAN	EST_HUMAN	NT
	Top Hit Acessian No.	(74870.1	(74870.1	(74870.1	4506608 NT	117795.1	4503044 NIT	4506728 NT	B028942 4	7152	4507152 NT	F193607.1	L163201.2	7879	0.0E+00 AA324262.1	Γ	4532	4504532 NT	4557887 INT	4557887 NT									8923955 NT	=373403.1	Г			
	Most Similar (Top) Hit BLAST E Value	0.0E+00)	0.0E+00.0	0.0E+001	0.0E+00	0.0E+00	0.01	0.0E+00	· 0.0E+00	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00	0.0E+00	0.0E+00	0.0E+00 B	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 AI	0.0E+00 AL163246.2	0.0E+00	0.0E+00 AB033035.1	0.0E+00 AU132898.1	0.0E+00 BE385144.1	0.0E+00 AW938825.1	0.0E+00 AL117233.1	0.0E+00	0.0E+00 BI	0.0E+00 AL163210.2	0.0E+00 BE081527.1	0.0E+00 BF028005.1	0.0E+00 A
	Expression Signal	3.82	3.87	3.87	114.34	1.06	2.62	36.55	3.49	6.44	6.44	3.81	0.76	3.88	1.31	2.44	5.86	5.86	21.4	21.4	2.88	5.35	5.35	3.11	1.82	4.41	4.79	1.51	0.81	0.68	4.2	1.9	1.86	1.25
	ORF SEQ ID NO:		10454			10059	10481		10482	10483	10484	10485		10498			10512	10513	10520	10521	10527	10528	10529	10534	10536	10542	10543	10545	10546		10555	10559	10565	10570
	Exon SEQ ID NO:						5463	5464	5465	5466	5466	5467	5478	5480	5485	5486	5502	5202	5506	5506	5517	5518	5518	925	228	5536	7722	5539	5540	5543	5550	7723	5562	5567
	Probe SEQ ID NO:	399	400	400	404	417	425	426	427	428	428	429	14	443	848	449	465	465	470	470	480	481	4	490	492	200	50	50	<u>8</u>	88	515	229	527	532

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Table 4
Single Exon Probes Expressed in HBL100 Cells

	_	<b>-</b>	-	_	_	_		_	_					_	_	_		_	_1	_	<u> </u>		4	<u>=</u>		4	1 _1		ą,	# 4.	9 4	<b>#</b>
Top Hit Descriptor	Homo sapiens transcription elongation factor B (SIII) polywanika 4 iika (TOEB41) - B414	Homo saciens guantine rucleotide binding profesh (Gradien) why 44 (Gradien) (Gradien)	Homo saplens guernne nucleotide binding protein (G protein) ainta 11 (Go class) (CNA141)	Homo saplens enillin (LOCS4443), mRNA	Homo septens emillin (LOCS4443), mRNA	Homo sepiens enillin (LOC54443), mRNA	Homo sepiens X-linked anhidrolitic ectodermal dyspiasta protein gene (EDA), exon 2 and flanking repeat redions	UI-H-BIT-acb-h-04-0-UI st NCI CGAP Sub3 Homo seniens cONA clare BAACE: 274-308-4-21	Homo sapiens RGH1 gene, retrovirus-fike element	Homo sapiens ubiquinol-cytochrome c reductase, Rieske iron-sulfur polypeptide 1 (UQCRFS1), nuclear gene ercoding mitochondrial protein, mRNA	Human abolipoprotein A-I (AppA-I) name emm 1	601822627F1 NIH MGC 75 Home septems china clone IMAGE-4045447 F.	Homo sapiens hypothetical protein FL/20701 (FL/20701), mRNA	Homo sapiens hypothetical protein FJ20701 (FL)20701), mRNA	Homo saplens hypothetical protein FLJ20701 (FLJ20701) mRNA	Homo sapiens hypothetical protein FLJ20701 (FLJ20701), mRNA	Homo sapiens hypothetical protein FLJ20701 (FLJ20701), mRNA	Homo sepiens hypothetical protein FLJ20701 (FLJ20701), mRNA	Homo sapiens Smad- and Olf-interacting zinc finger protein mRNA, partial cds	Homo seplens Smad- and Olf-interacting zinc finger protein mRNA, partial cds	Hamo sapiens NOD1 protein (NOD1) gene, exons 1, 2, and 3	Homo sapiens mRNA for KIAA 1386 protein, partial cds	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA	zt60c07.r1 Soares_testis_NHT Homo sepiens cDNA clone IMAGE:728732 57	Homo sapiens RGH2 gene, retrovirus-like element	Homo saplens novel SH2-containing protein 3 (NSP3) mRNA	Homo sapiens glutamate receptor, lonotropic, N-methyl D-aspartate 2B (GRIN2B) mRNA	Hamo saplens CCAAT-box-binding transcription factor (CBF2) mRNA	Human neutral amino acid transporter (ASCT1) gene, exon 8	Homo sapiens sodium/calcium exchanger isoform NaCa3 (NCX1) mRNA, complete cds
Top Hit Detabase Source	N-	보	N	N	NT	Z.	Ľ	T HUMAN	Г	— <b>5</b>	Į.	T HUMAN		Þ	Þ	5					NT	NT				EST_HUMAN						IN IN
Top Hit Acession No.	E006030 NT	4504038 NT	4504036 NT	8923831 NT	8923831 NT	8923831 NT	F003528.1	Γ	0.0E+00 D10083.1	5174742 NT	04066.1	0.0E+00 BF104898.1	23631	8923631 NT	8923631 NT	8923631 NT	8923631 NT	8923631 NT				0.0E+00 AB037807.1	6806918 NT	6806918 NT	6806918 NT	0.0E+00 AA399486.1		4885526 NT	6006003 NT	5031624 NT		-108389.1
Most Simitar (Top) Hit BLAST E Value			0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00/	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 /	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 U05235.1	0.0E+00 AI
Expression Signal	17.56	5.39	6:38	3.44	2.66	2.66	3.9	1.35	4.5	7.89	4.24	2.06	1.05	1.05	0.92	0.92	0.7	0.7	0.83	0.83	2.81	2.39	0.76	1.52	1.52	2.63	6.46	3.25	2.37	2.08	3.14	11
ORF SEQ ID NO:	10573	10574				10579		10588		10613	-	10627	10629	10630	10629	10630	10629	10630	10638	10639	10646	10649	10650	10651	10652	10662	10666		10678	10681	10685	10688
Exan SEQ ID NO:			5571		5574	5574	5578	5586	5595	5613	5625	5628	5630	5630	5630	5630	5630	5630	5637	5637	5644	5646	5648	5649	5649	5657	5661	5666	5673	5675	5678	5682
Probe SEQ ID NO:	535	536	536	538	539	539	644	552	261	581	594	287	599	539	909	009	691	601	610	910	617	619	621	622	622	629	633	638	645	947	920	654

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Hams saniens sodium/celcium exchanger isoform NeCe3 (NCX1) mRNA, complete ods	Light contents where X-linked (PROX) mRNA	Oliv cardina for the Control of the	Homo sapiens protean kinase, A-timaca (TransA) IIII u.k.a.	Human endogenous fettovirus pritt. 1 (ERVs)	Homo sapiens high-mobility group (nomistone chromosomal) process I (mino I) IIII was	Homo sapiens mRNA for KIAA1089 protein, partial cds	Homo sapiens similar to rat integral membrane glycoprotein POMIZI (POMIZILI), mixim	np49d01.s1 NCI_CGAP_B11.1 Hamo septens cDNA done IMAGE:1124033 3 snimer to go. No. 332. INTERFERON-INDUCIBLE PROTEIN 1-8U (HUMAN);	Human von Willebrand factor gene, exons 23 through 34	Human von Willebrand factor gene, exons 23 through 34	Homo sepiens TNF receptor-associated factor 1 (TRAF1) mRNA	Homo saplens ALR-like protein mRNA, partial cds	Hown carients ALR-like protein mRNA, partiel cds	Leave engines handhelited profess I 12(634 (FL)21634), mRNA	TO A DATO 770 Deciration on the metapopous leukernie cell (FAB M1) Baylor-HGSC project=TCAA Homo	CAAF 100 / 91 Charles action in your sealens cDNA clone TCAAP0779	Hown cariants MHC class I antiden (HLA-G) mRNA, HLA-G1 allele, complete cds	Homo sepiens MHC class   antigen (HLA-G) mRNA, HLA-G1 allele, complete cds	Luman electringues activator inhibitor-1 gene, exons 2 to 9	Fluitian, plantimoses activator britishor-1 gene, exons 2 to 9	Homo seniens mRNA for KIAA1339 protein, partial cds	Homo gapiens zinc finder protein 212 (ZNF212), mRNA	Homo sapiens mRNA for repressor protein, partial cds	601445647F1 NIH MGC 65 Home septens cDNA clone IMAGE:3849803 5	vicson08 r1 Soures breast 2NbHBst Homo sepiens cONA clone IMAGE:154046 6	Homo saplens splicing factor 3a, subunit 1, 120kD (SF3A1), mRNA	Homo sablens derre for AF-8, complete cds	Homo sepiens KIAA0170 gene product (KIAA0170), mRNA	Human mRNA for KIAA0184 gene, pertial cds	Himman mRNA for KIAA0184 gene, partial cds	H seniens mRNA for Interferon alpha/beta receptor (long form)	Homo sapiens mRNA for KIAA0910 protein, partial cds	Homo saplens mRNA for KIAA0910 protein, partial cds	
	Top Hit Database Source									EST HUMAN								- For	NOMO!		-	_	-15	-	2 2	OT LINKIAN	EST JUNAN	אים שרוש ביי	-	I E	ž l	= +	Z L	2 12	12	
		-		Ę	된	IN	N A	E	2	-83	호	Y	Z	1	4	-	킭	ŭ	3	4		z			Z Z 2	+	1	9		2			+	Ť	1	1
	Top Hit Acession No.		10836	4826947 IN	4826947 NT	X57147.1	4504424 NT	0.0E+00 AB029012.1	7657468 NT	0 0F+00 AA614537.1	0.0E+00 M60675.1	0 0F+00 M60875 1	TN 29192 NT	A CO64750 4	0.0E+00 AF 264750.1	0.0E+00 AF264750.1	11545800INI		0.0E+00 BE2415//.1	0.0E+00 AF225990.2	0.0E+00 AF226990.2			AB03//6				K48915.1	_	AB01139	_+	_	_	-	AB020717.1	JABUZUI II.I
	Most Similar (Top) Hit BLAST E Value		0.0E+00 Ar	0.0E+00	0.0E+00	0.0E+00 X57147.1	0.0E+00	0.05+00	00±400	0.05+00	0.0E+00	00+100	200	0.05.00	0.0=+00	0.0E+00	0.0E+00		0.0=+00	1						١	-	1						١	١	0.05+00
	Expression Signal		=	4.21	4.21	1.79	21 88	45.74	289	17.08	404	70,	4.04	1.32	3.44	3.44	11.53		2.72	1.78	1.78	3.03		1.73												3.13
	ORF SEQ ID NO:	-	10689	10694	10695		40708				10738					10760	10763		10771	10795	10796	10799		10801		10804		10809	10810	1 10819		5 10835	5 10836	0 10840		10845
	Exan SEQ ID NO:		5682	2887	5687		L	1			5727					5742	5744		6750	5769	5769	L	5772	2 5774		5 7730	5777	5781	1 5782	0 5791	3 5795	5805		9 5810	5814	3 5814
	Probe SEQ ID NO:		654	099	8	988	3	0/4	8/8	880	989	3	733	713	719	719	721		727	747	147	750	750	762	753	75	75	186	79	230	773	787	784	789	793	793

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Single Exon Probes Expressed in HBL100 Cells

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Human ras inhibitor mRNA, 3' end	Human ras inhibitor mRNA, 3' end	Human ras inhibitor mRNA, 3' end	Homo sepiens thyrotrophic embryonic factor (TEF), mRNA	Homo saplens thyrotrophic embryonic factor (TEF), mRNA	0598e03.s1 NCI_CGAP_GC3 Hamo saplens cDNA done IMAGE:16134043'	os98e03.s1 NCI_CGAP_GC3 Homo sepiens cDNA done IMAGE:1613404 3'	Homo sepiens KIAA0929 protein Msv2 Interacting nuclear farget (MINT) homolog (KIAA0929) mRNA	Homo sepiens mRNA for PSP24, complete cds	PM2-GN0014-050900-001-f02 GN0014 Homo sapiens cDNA	PM2-GN0014-050900-001-f02 GN0014 Homo seplens cDNA	PM2-GN0014-050900-001-f02 GN0014 Homo saplens cDNA	Homo sapiens partial C-fgr gene, exons 2 and 3	Homo sapiens partial c-fgr gene, exons 2 and 3	Homo sapiens chromodomain protein, Y chromosome-like (CDVL) mRNA	Human beta-tubulin (TUB4q) gene, complete cds	Human beta-tubulin (TUB4q) gene, complete cds	Human beta-tubulin (TUB4q) gene, complete cds	Homo sapiens 8q22.1 region and MTG8 (CBFA2T1) gene, partial cds	Homo sapiens 8q22.1 region and MTG8 (CBFA2T1) gene, pertial cds	Homo sapiens 14q32 Jagged2 gene, complete cds; and unknown gene	Homo sapiens 14q32 Jagged2 gene, complete cds; and unknown gene	Homo sapiens 14q32 Jagged2 gene, complete cds; and unknown gene	Homo sapiens 14q32 Jagged2 gene, complete cds; and unknown gene	Homo sapiens DKFZP586M0122 protein (DKFZP586M0122), mRNA	Homo sapiens Inner membrane protein, mitochondrial (mitofilin) (IMMT), mRNA	aa86g07.s1 Stratagene fetal retina 837202 Homo saplens cDNA clone IMAGE:838236 3' similar to	SW.PRS8_HUMAN P47210 28S PROTEASE REGULATORY SUBUNIT 8;	EST51124 WATM1 Hamo septens cDNA clane 51124 similar to DNA-DIRECTED RNA POLYMERASE II	(alignment Ser and Pro with BLASTx or p)	EST61124 WATM1 Home sapiens CDNA clone 61124 similar to DNA-DIRECTED RNA POLYMERASE II	(alignment Set and Prowith BLASIx or p)	Homo sapiens TRAF family member-associated NFKB activator (TANK) mRNA
Top Hit Database Source	LN	L	TN.	Z	Ę	EST HUMAN	EST_HUMAN	_ <u></u>	Ę	EST HUMAN	EST HUMAN	EST_HUMAN	LZ.	N.	LN.	TN	E	ΤŃ	NT	NT	NT	NT	٦N	LN	FN	FZ		EST_HUMAN		EST_HUMAN		EST_HUMAN	¥
Top Hit Acession No.		M37190.1	M37190.1	4507430 NT	4507430 NT	AI001948.1	A1001948.1	7657268 NT	AB030566.1	3F366974.1	3F366974.1	3F366974.1	X52207.1	X52207.1	4757969 NT	J83668.1	J83668.1	J83668.1	4F198490.1			4F111170.3		4F111170.3	7681685 NT	5803114 NT		4A458680.1		V43182.1		V43182.1	4759249
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00/		0.0E+00	i i	0.0E+00	0.0E+00
Expression Signal	1.21	6.98	0.78	1.55	1.55	1.94	1.94	7.69	1.89	1.14	1.14	1.14	1.55	1.55	2.93	3.09	58.93	27.22	5.92	11.99	1.12	1.89	1.74	6.04	2.62	4.81		4.43		0.8		8.0	1.08
ORF SEQ ID NO:	10985	10986	10987				10097	10999	11009	11015	11016	11017	11018	11019	11026	11034	11035	11035			11040	11040	11040	11041	11044	11048				11052		11053	11054
Exan SEQ ID NO:		٠	5955			7735	7735	5962	5975	5983	5983	5983	5984	5984	5993	6003	6004	6004	6007	6007	6010	6010	6010	6011	6014	6018		6019		6022		6022	6023
Probe SEQ ID NO:	936	837	938	939	839	947	947	949	096	896	896	896	696	696	978	686	066	991	994	995	866	666	1000	1001	1004	1008		1009		1012	,	1012	1013

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo septens TRAF family member-associated NFKB activator (TANK) mRNA	Homo saplens hypothetical protein PL 11196 (FL 11196), mRNA	Homo sapiens heat shock 70kD protein 98 (mortelin-2) (HSPA98) mRNA	Homo seplens cedherin 6. K-cadherin (fetal kidney) (CDH6) mRNA	Homo sapiens cadherin 6, K-cadherin (fetal kidney) (CDH6) mRNA	Homo saplens hypothetical protein FL/20695 (FL/20695), mRNA	Homo sapiens hypothetical protein FLJ20695 (FLJ20695), mRNA	Homo sapiens mRNA for alpha-tubulin 8 (TUBA8 gene)	Homo sapiens hypothetical protein FLJ20080 (FLJ20080), mRNA	Hamo sapiens alkylation repair; alkB hamolog (ABH), mRNA	Homo sapiens Death associated protein 3 (DAP3) mRNA	MR0-BN0115-200300-003-h08 BN0115 Homo sapiens cDNA	Homo sapiens potassium channel, subfamily K, member 9 (KCNK9), mRNA	Homo sapiens potassium channel, subfamily K. member 9 (KCNK9), mRNA	Homo sapiens protein kinase, X-finked (PRKX) mRNA	Homo sapiens protein kinase, X-linked (PRKX) mRNA	Homo sapiens ribosomal protein S27a (RPS27A) mRNA	Homo sapiens hypothetical protein FLJ20309 (FLJ20309), mRNA	Homo sapiens DNA for Human P2XM, complete cds	Homo sepiens DNA for Human P2XM, complete cds	Homo sapiens similar to rat integral membrane glycoprotein POM121 (POM121L1), mRNA	Homo sapiens similar to rat integral membrane glycoprotein POM121 (POM121L1), mRNA	Homo sapiens Npw38-binding protein Npw8P (LOC51729), mRNA	H.sapiens ART4 gene	H.sapiens ART4 gene	qb22d10.x1 Soares_pregnant_uterus_NbHPU Homo sapiens cDNA clone IMAGE:1697011 3'	Homo sapiens mRNA for KIAA0903 protein, partial cds	Homo sapiens chondroitin sulfate proteoglycan 2 (versican) (CSPG2) mRNA	Homo sapiens chondroltin sulfate proteoglycan 2 (versican) (CSPG2) mRNA	Homo sapiens chromosome 12 open reading frame 3 (C12ORF3), mRNA	Homo sapiens glutamate decarboxylase 1 (brain, 67kD) (GAD1), transcript variant GAD25, mRNA	Homo sapiens glutamate decarboxylase 1 (brain, 67kD) (GAD1), transcript variant GAD25, mRNA	Homo saplens mRNA for KIAA1414 protein, pertial cds	Homo saplens keratin 18 (KRT18) mRNA
	Top Hit Database Source	Ę	7	L	TN.	٦	LZ.	Z					EST_HUMAN							NT	12						EST_HUMAN	LN							
6	Top Hit Acession No.	4759249 NT	8922933 NT	4758569 NT	4826672 NT	4826672 NT	8923624 NT	8923624 NT	J245922.1	TN 8923087	5174384 NT	4758117 NT	E005208.1	7706134 NT	7706134 NT	4826947 NT	4826947 NT	4506712 NT	8923290 NT	B002059.1	\B002059.1	7657468 NT	7657468 NT	7706500 NT	(95826.1		1147650.1	.B020710.1	4758081 NT	4758081 NT	9965844 NT	7305076 NT	5076	B037835.1	4557887 NT
-	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 B	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	$\sim$	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00
	Expression Signal	1.08	5.63	10	3.04	3.04	3.86	3.86	127.96	1.04	4.28	3.16	5.76	4.27	4.27	1.67	1.67	46.69	2.22	37.23	82.18	4.76	4.76	4.36	0.66	99.0	1.23	3.06	0.86	0.86	1.05	5.48	5.48	0.98	14.52
	ORF SEQ ID NO:	11055		11070	11085	11086	11090	11091	11092		11096	11107	11115	11139	11140	11150	11151	11152	11154	11157	11158	11159	11160	11162	11163	11164	11165	11167	11173	11174	11175	11186	11187	11189	11198
	Exan SEQ ID NO:	6023	6026	6040	2909	6057	6061	6061	6062	6064	9909	6074	6086	6109	6109	6121	6121	6122	6124	6127	6129	6130	6130	6132	6133	6133	6134	6136	6143	6143	6144	6154	6154	6156	6163
	Probe SEQ ID NO:	1013	1016	1030	1048	1048	1052	1052	1053	1055	1057	1066	1079	1102	1102	1115	1115	1116	1118	1121	1123	1124	1124	1127	1128	1128	1129	1131	1138	1138	1139	1150	1150	1152	1159

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Homo sapiens Na+fil+ exchancer (softom 2 (NHF2) mRNA complete ode	Homo saciens mut. (E. coll) hamafar 3 (All H3) mRNA	Homo sepiens hypothetical protein FL/10697 (FL/10697) mRNA	Homo saplens ALR-like protein mRNA partial ods	Homo saplens ALR-like protein mRNA, partial cds	Homo sapiens ALR-like protein mRNA, partial cys	Homo sapiens ALR-like protein mRNA, pertial cds	Homo sapiens chromosome 3 subtelomeric region	Homo saplens chondroitin sulfata profesoriwan 4 (melanoma-associated) (CSBCA)	Homo saciens prefidin 4 (PEDNA) mRNA	Homo saciens NF2 cene	Homo saplens ribosomal protein S2 (RPS2) mRNA	Homo sapiens Williams-Beuren syndrome deletion trenscorint o (WIRCODO) months	Homo saplens mRNA for KIAA1507 barden naring ode	Homo sapiens mRNA for KIAA1507 protein pertial cds	Homo sapiens Wolfram syndrome (WFS) mRNA	Homo saplens Wolfram syndrome (WFS) mRNA	Homo sapiens Woffram syndrome (WFS) mRNA	Homo saplens protein phosphatase 2A BR gamma subunit gene earn 5	Homo sepiens rhebdoid tumor deletion region protein 1 (RTDR1), mRNA	Homo sapiens rhabdoid turnor deletion region protein 1 (RTDR1), mRNA	Homo sapiens ring finger protein 9 (RNF9), mRNA	Homo saplens zinc finger protein 173 (ZNF173) mRNA	Homo sapiens ring finger protein 9 (RNF9), mRNA	Homo saplens zinc finger protein 173 (ZNF173) mRNA	Homo sapiens mRNA for KIAA0577 protein, complete cds	Homo sapiens KIAA0170 gene product (KIAA0170), mRNA	Homo sepiens KIAA0170 gene product (KIAA0170), mRNA	Hamo sepiens period (Drosaphilia) hamolog 3 (PER3), mRNA	Homo sapiens period (Drosophila) homolog 3 (PER3), mRNA	Human endogenous retrovirus HERV-K10	601109792F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3350471 5'	601109792F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3350471 5'	Homo sapiens mRNA for Familial Cylindromatosis cyld gene
Top Hit Database Source	N	5	5	IN	Į.	۲	LN TA	IN				1							Į.												- LV		T_HUMAN	- L
Top Hit Acession No.	0.0E+00 AF073299.1	7857336 NT	8922593 NT	26475	264750.1	264750.1	264750.1	109718.1	4503098 NT	4505740 NT	718000.1	4508718	0.0E+00 AF084479.1	Γ		5174748 NT	5174748 NT	5174748 NT	0.0E+00 AF096156.1	7657529 NT	7657529 NT	5803146 NT	4508004 NT	5803146 NT	4508004 NT	0.0E+00 AB011149.1	7661965 NT	7661965 NT	8567387 NT	8567387 NT	4123.1			0.0E+00 AJ250014.1
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00 AF	0.0E+00 AF	0.0E+00 AF	0.0E+00 AF	0.0E+00	0.0E+00	0.0E+00	0.0E+00 Y18000.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.05+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 M1	0.0E+00	0.0E+00	0.0E+00/
Expression Signal	1.01	1.32	0.69	0.89	0.89	1.55	1.09	4.89	2.52	4.31	2.72	283.18	5.64	1.42	1.42	2.45	2.45	2.45	2.35	1.16	1.16	1.37	2.26	0.67	2.07	4.17	9.28	10.18	3.67	3.67	1.22	1.64	1.64	1.51
ORF SEQ ID NO:	11209		11241	11244	11245	11246	11247	11272	11273	11279		11294	11303	11307	11308	11321	11322	11323		11335	11336	11341	11342	11343	11344	11346	11347	11348	11349	11350	11363	11421	11422	11431
Exan SEQ ID NO:										6236	6245	6253	6260	6266	6266	6280	6280	6280	6281	7743	7743	6294	6295	6297	6298	8300	33	6302	6303	6303	6314	6372	6372	6381
Probe SEQ ID NO:	1172	1190	1203	1206	1206	1207	1208	1227	1228	1238	1247	1255	1262	1268	1268	1281	1281	1281	1282	1292	1292	1296	1297	1239	1300	1302	1303	1304	1305	1305	1317	1375	1375	1384

Page 183 of 209 Table 4 Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	qg38b08.x1 Soares_testis_NHT Homo sepiens cDNA clone MAGE:1837427 3' simiter to WF:12/A1.5 CE14213;	RAN, member RAS oncogene familyHomo sepiens RAN, member RAS oncogene (emity (RAN), mRNA	Homo sapiens proprotain convertase subditts in Kedin type Z (PCSNZ) many	Homo sapiens proprofein convertase subtilism/Kedin type Z (FCSNZ) minus	Homo saplens KIAA1114 protein (NIAA1114), minna	Homo sapiens KiAA1114 protein (NIAA1114), tilinum	Homo sapients partial Art 4 gents, auchs z to 7 and 1 chorus communications are all the second and the second and the second are all the second and the second and the second and the second are all the second and the	Homo saprats aprilla i cucco ju anistatato permit ci i i i i i i i i i i i i i i i i i	Nove numan general on university as	Nove numan gene mapping to drontoeding	Human mRNA for KIAAU240 gene, parum cus	Homo sapiens calcineum binding protein 1 (NICANASSA), IIII AAA	Homo saplens KIAA0170 gene product (KIAA0170), minner	Homo sapiens KIAAU170 gene product (Nirvo 1707, minute)	Home sapiens made to nonered of Dieselphina Institute Colors (1000)	EST371757 MAGE resequences, MAGE noting septems control of the State o	Bastals, TNCI_COAT_CODI_INIDERCATE TO THE AMERICAN COLOR TO THE AM	Cercopitheous aethiops cyclophillin A mixiva, complete cds	Cercopinedus acumpts cyclopinimi A ministri compressiones de la compressione de la compre	ESTSSSON MACH resentences, MAGN Homo septens CDNA	EST SOSSO WAY C. Conference of the sosso of	Louis contours Britan's transing kinase (BTK), alpha-D-galactosidase A (GLA), L44-like ribosomal protein	(L44L) and FTP3 (FTP3) genes, complete cds	Homo sapiens transmembrane glycoprotein (GPNMB) mRNA	Homo sapiens transmembrane glycoprotein (GPNMB) mtkNA	Homo sapiens KIAA0957 protein (KIAA0957), mRNA	Homo sapiens TNF-inducible protein CG12-1 (CG12-1), mRNA	Human trensclutzminese mRNA, complete cds	Homo saciens tith (TTN) mRNA	Homo sapiens tith (TTN) mRNA	Homo saplens ribosomal protein L5 (RPL5) mRNA	
-	Top Hit Database Source	EST_HUMAN				1=						7		1	<u></u>	_	EST_HUMAN	EST HUMAN	<b>L</b>	<b>-</b>	EST HUMAN	ESI HUMAN		<del></del>	٥	_	1		H		<u> </u>	= =====================================	
-		S	<u> </u>	N N	N S	<u>   </u>	<u>S</u>	킬	Ż	킼	뉟	Ż	57 N	N 99	S5 N	34 N	ü	ŭ	뉟	Ż		7	=	_ <u>Z</u>	4505404 N	Š	S	7656072 NT			TA 007/004	450//20 NI	5
	Top Hit Acession No. ·	1208756.1	6042206 NT	4505846 NT	4505846 NT	7705565 NT	7705565 NT	4,7238093.1	4F038280.1	AL132999.1	AL137764.1	D87077.1	6912457 NT	7661965 NT	7661965 NT	7706434 NT	AW959687.1	AA4811721	AF023860.1	AF023860.1	AW976097.1	AW976097.1	D10884.1	U78027.1						M984/8.			
	Most Similar (Top) Hit BLAST E Value	0.0E+00 A	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	000+000	00+30 o	1						١	0.0E+00
	Expression Signal	1.16	28.34	1.59	1.59	3.9	3.9	4.32	3.56	2.12	1.3	1.69	6.31	2.08		3.1		2.99	42.31	42.31			1.11	4.87					4.7				39.26
	ORF SEQ ID NO:	11442	11443	11453	11454	_			11470	11487	11488	L	}_	L		L	L				11555	11556	11557					11562				4 11571	
	Exan SEQ ID NO:	8389	San A	8388	6398	9400	6400	6402	6411	6431	١.					L			L			6500	6501		1	_			3 6506	3 6511	5 6514		7 7749
	Probe SEQ ID NO:	1392	293	1404	169	1403	1403	1405	1413	1434	1435	1439	4442	1444	1444	4480	1493	4404	1500	1500	1502	1502	1503		1505	1506	1506	1507	1508	1513	1516	1516	1517

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Table 4
Single Exon Probes Expressed in HBL100 Cells

																				p	مسة	. 4 .	- 4				-		- 4	-	4	بساه ج
Top Hit Descriptor	Human laminin receptor (2H5 epitope) mRNA, 5' and	Homo saplens chondrolitin sulfate proteoglycan 4 (melanome-associated) (CSPCA) mRNA	human o-yes-2 gene	H. sapiens hH2B/e gene	Homo sapiens butyrophilin, subfamily 2, member A1 (BTN2A1), mRNA	Homo saplens butyrophilin, subfamily 2, member A1 (BTN2A1), mRNA	AV690831 GKC Homo sepiens cDNA clone GKCBOF02 5	AV690831 GKC Hamo saplens cDNA clane GKCBOF02 5	Homo sapiens mRNA for KIAA1472 protein, partial cds	Homo sapiens DNA polymerase zela catalytic subunit (REV3) mRNA, complete cds	Homo sapiens KIAA0569 gene product (KIAA0569), mRNA	Homo sapiens KIAA0569 gene product (KIAA0569), mRNA	Homo sepiens heat shock 70kD protein 10 (HSC71) (HSPA10), mRNA	Homo sapiens heat shock 70kD protein 10 (HSC71) (HSPA10), mRNA	Human sodium channel mRNA	yo76c05.s1 Soares adult brain N2b4HB55Y Homo saplens cDNA clone IMAGE:183848 3'	Homo sapiens mRNA for KIAA1609 protein, partial ods	Homo sapiens mRNA for KIAA1609 protein, partial cds	UI-H-Bi3-ejw-<-04-0-UI.s1 NCI_CGAP_Sub5 Homo septens cDNA clone IMAGE:2733284 3'	MR0-HT0166-191199-004-b11 HT0166 Homo sepiens cDNA		wg81b07.x1 Soares_NSF_F8_9W_OT_PA_P_S1 Homo sepiens cDNA clone IMAGE:2371477 3' similar to TR:062788 062788 CYS2HIS2 ZINC FINGER PROTEIN	ZNF1) mRNA		pods	Human zinc-finger protein 7 (ZFP7) mRNA, complete cds	Homo sapiens keratin 18 (KRT18) mRNA	Homo saplens vets avian erythroblastosis virus E26 oncogene related (ERG), mRNA	hu11d05.x1 NCI_CGAP_Lu24 Homo eaplens cDNA clone IMAGE:3166281 3' similar to TR:095147 095147	MACH LIKE THO I EIN I TROSINE PHOSPHATASE:	hu11d05.x1 NCI_CGAP_Lu24 Homo sepiens cDNA clone IMAGE:3166281 3' similar to TR:095147 095147 MKP-1 LIKE PROTEIN TYROSINE PHOSPHATASE ;	Homo sapiens gamma-aminobutyric acid (GABA) A receptor, gamma 2 (GABRG2) mRNA
Top Hit Database Source	¥	Z.	F	Z	N.	F	EST_HUMAN	EST HUMAN	L L	LN.	N F	N.	Ę	N.	Z	EST_HUMAN	NT	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	N-I	Į.	Z	LN	Z	T'A		EST HUMAN	EST_HUMAN	LZ.
Top Hit Acession No.	M14199.1	4503098 NT	D00333.1	283738.1	5921460 NT	5921460 NT	AV690831.1	AV690831.1	AB040905.1	AF157478.1	7662183 NT	7662183 N.T	5729876 NT	5729876 NT	M91803.1	H26973.1	AB046829.1		AW 444637.1	BE144364.1	BE144364.1	Al768104.1	4758513 NT	AF057177.1		M29580.1	4657887 NT	7657065 NT	7 7 20000	BE2223/4.1	BE222374.1	57610
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00 H	_		0.0E+00	_	0.0E+00 E	0.05+00/	_	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0=+00	0.0E+00	0.0E+00
Expression Signal	61.77	. 8.55	1.85	26.28	2.84	2.84	6.07	6.07	1.72	2.77	6.44	6.44	84.9	84.9	2.3	9.86	1.95	1.95	4.22	8.38	8.38	3.34	1.18	2.39	2.07	207	28.94	1.45	07.0	2.18	2.18	1.20
ORF SEQ ID NO:	11572	11587		11601	11602	11603	11604		11608			11613	11614	11615	11617	11631	11639	11840	11655	11690	11691	11695	11696	11697	11700	11701	11703	11704	202777	11/0/	11708	11710
H H H	6515	6528	6535	6543	6544	6544	6545	6545	7750	6550	6552	6552	6554	6554	6556	6269	6576	6576	6594	6622	6622	6626	6627	6628	6631	6631	6633	6634	0000	88	8638	6839
Probe SEQ ID NO:	1518	1530	1537	1545	1546	1546	1547	1547	1549	1553	1555	1555	1557	1557	1559	1572	1579	1579	1598	1625	1625	1629	1630	1631	1634	1634	1636	1637	4014	104	1641	1643

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Table 4
Single Exon Probes Expressed in HBL100 Cells

			_	_	-	_	_	_		_	-	_	_	_	_	_	_	H	-		بية °د	* ***	****		-	_	*****		_
Top Hit Descriptor	1059608.rt Source breest 3Nbi-HBst Homo explens cDNA clone IMAGE:182246 6' similar to gb:M64099 GAMMA-GLUTAMM_TRANSPEPTIDASE 5 PRECURSOR (HLIMAN):	y59908.rl Scares breast 3Nb/HBs Homo eaplens cDNA clone MAGE:182246 6' similar to gb:M64099 GAMMA-GLUTAAM, TRANSPEPTIONSE R PRECY IDSOND KIN MANNE.	H. sapiers H.28 h dene	H. sapiens H2B/h gene	Homo saplens high-mobility group (nonhistone chromosomal) protein 17 (HAACAT)	Homo sapiens FOXJ2 forkhead factor (LOCSS610) mRNA	Homo sapiens pericentriolar material 1 (PCM1) mRNA	Homo saplens RNA binding motif protein. Y chromosome, family 1, member A1 (RBM/V1A1) mRNA	Homo sepiens WAVE2 mRNA for WASP-family protein, complete cds	TCR zeta [human, Genomic/mRNA, 365 nt, segment 1 of 8]	Homo sapiens solute carrier family 26 (sulfate transporter), member 2 (SLC26A2) mRNA	Homo sapiens NOD2 protein (NOD2), mRNA	Homo sapiens SMCY (SMCY) gene, complete cds	Homo sapiens ribosomal protein S2 (RPS2) mRNA	Homo sapiens E1A binding protein p300 (EP300) mRNA	Homo saplens E1A binding protein p300 (EP300) mRNA	Human CSF-1 receptor (FMS) gene, complete cds, and (SMF) gene, pertial cds	Homo sapiens nuclear autoantigenic sperm protein (histona-binding) (NASP) mRNA	Human ribosomal protein L21 mRNA, complete cds	Human mRNA for KIAA0333 gene, partial cds	Homo septens activating transcription factor 4 (tax-responsive enhancer element B67) (ATF4) mRNA	Homo sapiens activating transcription factor 4 (tax-responsive enhancer element B67) (ATF4) mRNA	Homo sepiens activating transcription factor 4 (tax-responsive enhancer element B97) (ATF4) mRNA	Homo saplens immunoglobin superfamily, member 3 (IGSF3) mRNA, and translated products	Homo sepiens immunoglobin superfamily, member 3 (IGSF3) mRNA, and translated products	Homo sapiens Retina-derived POU-domain factor-1 (RPF-1), mRNA	Homo sapiens Retina-derived POU-domain (actor-1 (RPF-1), mRNA	Homo sapiens mRNA for KIAA1152 protein, partial cds	Homo sapiens mRNA for KIAA1152 protein, partial cds
Top Hit Detabase Source	EST_HUMAN	EST HUMAN	Z	Z,	N.	F	Ę	F	5	FZ	N <sub>z</sub>	Ę	NT	NT	N-	N F	Z	NT	Z	N1	-N	_ <u>F</u>	_ <del>\</del>	NT.	Ę	Ę	NT		L
Top Hit Acession No.	H30132.1	H30132.1	Z80780.1	280780.1	5031748 NT	8923841 NT	5453855 NT	4826973 NT	AB026542.1	S94400.1	4557538 NT	11545911 NT	AF273841.1	4506718 NT	4557558 NT	4557556 NT	U63963.1	4505332 NT	U14967.1	AB002331.1	4502264 NT	4502284 NT	4502284 NT	4504626 NT	4504626 NT	6005855 NT	6005855 NT		4B032978.1
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
Expression Signal	8.67	8.67	19.26	19.28	64.75	5.23	1.85	1.1	7.62	1.82	1.82	1.05	2.59	150.64	3.32	3.32	2.75	8.02	22.06	12.66	13.86	13.86	13.86	1.29	1.29	5.58	5.58	3.27	3.27
ORF SEQ ID NO:	11713	11714	11716	11717		11730	11733	11741	11748		11757	11764	11778		11826	11827	11831	11836	11851	11854	11855	11856	11857	11872	11873	11878	11879	11888	11889
Exam SEQ ID NO:	6642	6642	6844	6644	6647	6656	6659	8665		6673	6682	7753	6701	7754	6746	6746	6748	7755	6763	6765	6766	6766	8766	6780	6780	6289	62/8	6798	6798
Probe SEQ ID NO:	1646	1646	1648	1648	1651	1660	1663	1669	1675	1677	1686	1693	1706	1747	1752	1752	1755	1759	1771	1773	1774	1774	1774	1788	1788	1798	1798	1807	1807

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Table 4
Single Exon Probes Expressed in HBL100 Cells

		F	_	_	_		_	_	_			_	_	_		_		-	4		<u> </u>		1		1	_0_	<u> </u>	4	Ų.,,	# C	<b>3</b> 40
Top Hit Descriptor	Homo septens potassium voltage-gated channel, Shab-related subfamily, member 1 (KCNB1) mRNA	Homo sandens nodessalm valiane and choose should sale as the sale of the sale	Human retinal decementation slow (RDS) own Amon 1	Human retinal deceneration slow (RDS) cana awn 1	UI-H-BIT-effn-f-07-0-UI-st NCI CGAP Sub3 Home senions CPNA claus MAACE: 07202000 01	UI-H-B(1-efn-f-07-0-UI.s1 NCI CGAP Sub3 Homo seniers cDNA clara MACF-27273333	601179164F1 NIH MGC 20 Homo sapiens DNA clone IMAGE:3547279 51	601179164F1 NIH_MGC_20 Homo saplens cDNA clone IMAGE-3547239 F	Homo sapiens nuclear protein (NP220), mRNA	Homo saplens nuclear protein (NP220), mRNA	Homo sapiens RAD1 (S. pombe) homolog (RAD1) mRNA, and translated products	Homo sapiens RAD1 (S. pombe) homoloo (RAD1) mRNA and translated provisites	Homo sapiens mRNA for KIAA1367 protein, pertiel cds	Homo saplens DNA polymerase zeta catalytic subunit (REV3) mRNA, complete cds	Human transglutaminase mRNA, complete cds	Human transglutaminase mRNA, complete cds	Homo sapiens transforming growth factor, beta 3 (TGFB3), mRNA	Homo seplens transforming growth factor, beta 3 (TGFB3), mRNA		Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1) cones. Complete cds	e   pseudogene 1	, member A2 (BTN3A2), mRNA	5				MRNA				
Top Hit Database Source	Ę	<u> </u>	NT TA	Į.	EST HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	K	FZ	Z	N	N.	LN L	NT	TN	NT	Ä	LN T	- 4	NT	LN.	EST_HUMAN	N	۲	TN.	NT	TN	NT		EST_HUMAN
Top Hit Acession No.	4826783 NT	4826783 NT	U07147.1	J07147.1	AW207280.1	4W207280.1	3E277465.1	3E277465.1	TN 0657390	7657390 NT	4506384 NT	4506384 NT	B037788.1	4F157476.1	198478.1	198478.1	4507464 NT	4507464 NT	7657038 NT	0.0E+00 AF240788.1	A55632.1	5901905 NT	3E018066.1	4809282 NT	4809282 NT	4826638 NT	4826638 NT				
Most Similar (Top) Hit BLAST E Vatue	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 B	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 M33782.1	0.0E+00 M33782.1	0.0E+00 /	0.0E+00/
Expression Signal	3.31	3.31	8.06	8.06	1.53	1.53	2.87	2.87	1.79	1.79	2.53	2.53	1.87	1.55	1.49	1.49	1.31	1.31	1.17	6.49	3.82	1.74	4.05	1.47	1.47	9.6	9.2	1.41	1.41	1.95	1.95
ORF SEQ ID NO:	11891	11892	11893	11894			11915	11916	11955	11956	11958	11959	11967		11970	11971	11978	11979	11981			11988	11990	11995	11996	12007	12008	12023	12024	12025	12026
Exan SEQ ID NO:	6801	6801	6802				6827			6867	6870	6870	6877	6880	7758	7758	6885	6885	6887	6889	6894		9889	6902	6902	6912	6912	6926	8269	6928	6928
Probe SEQ ID NO:	1811	1811	1812	1812	1815	1815	1837	1837	1878	1878	188	1881	1888	189	1892	1892	1897	1897	198	1902	1907	1908	1910	1916	1916	1926	1926	1940	1940	1942	1942

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Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Home saplens calcinaurin binding protein 1 (KIAA0330)	Homo sapiens celcheum binding protein 1 (Kitahasa)DXIA	Homo sapiens KIAA0408 gene product (KIAA0408) mBNA	Homo spoiens mRNA for KIAA0577 british complete ade	H. sapiens genes for semenacelin I and semanacelin II	H.sapiens genes for semenocelin I and semenorelin II	Homo saplens mRNA for KIAA1513 protein, partial cds	Homo sapiens SMCY (SMCY) oene, complete cds	Homo sapiens SMCY (SMCY) oene complete ods	Homo sepiens TP53TG3e (TP53TG3e) mRNA	601573895F1 NIH MGC 9 Homo saciens of NA clear MACE: RRYETOR R	601573895F1 NIH MGC 9 Hamp sapiens cDNA clane MAGE:3835198 F1	Homo saplens coagulation factor IX (plasma thrombopiastic component, Christmas disease, hemophilia B) (F9) mRNA	601861974F1 NIH MGC 53 Homo sapiens cDNA clone IMAGE-40R1483 F	AU140831 PLACE4 Homo sapiens cDNA clone PLACE4000321 5'	7B22E10 Chromosome 7 Fetal Brain cDNA Library Homo seniess cPNA close 7B22E10	7822E10 Chromosome 7 Fetal Brain cDNA Library Homo saniens china choa 7822E10	Homo saplens similar to rat integral membrane alycoprotein POM/121 (POM/1211.1) mRNA	Homo sapiens phosphodiesterase 6A, cGMP-specific, rod, alpha (PDF6A) mRNA	HSCOIC021 normalized infant brain cDNA Homo sapiens cDNA close c-0ic02	qv90f08.x1 NCI_CGAP_UI2 Homo septens cDNA clone IMAGE:1988871.3' similar to contains Alu repetitive element;	601485148F1 NIH MGC 69 Homo sapiens cDNA clone IMAGE:3887747 5	601902604F1 NIH_MGC_19 Homo sepiens cDNA clone IMAGE:4135320 57	601902604F1 NIH_MGC_19 Homo sepiens cDNA clone IMAGE:4135320 57	RC3-CT0413-270700-022-d10 CT0413 Homo sapiens cDNA	RC3-CT0413-270700-022-d10 CT0413 Homo sepiens cDNA	Human plasma membrans calcium ATPase Isoform 2 (APT2B2) mRNA, comiete cds	Human plasma membrane calcium ATPase isoform 2 (APT2B2) mRNA, comlete cds	Homo saplens GTP binding protein 1 (GTPBP1) mRNA	QV1-GN0065-140800-318-c10 GN0065 Homo sapiens cDNA	Homo sapiens X-linked Juvenile retinoschisis protein (XLRS1) gene, exon 6 and complete cds	601672066F1 NIH MGC 20 Home sapiens cDNA done IMAGE:3954785 5
	Top Hit Detabase Source	Z	NT.	LN LN	F	Z	N F	K	N	Į.	LN	EST HUMAN	EST_HUMAN	   <u> </u>	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	L	N	EST_HUMAN	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	NT	IN	77	EST HUMAN	L'A	EST_HUMAN
	Top Hit Acession No.	6912457	6912457 NT	7662095 NT	AB011149.1	247556.1	247556.1	4B040946.1	4F273841.1	AF273841.1	7706742 NT	3E743215.1	E743215.1	4503648 NT	F207688.1	0.0E+00 AU140831.1	0.0E+00 AA077589.1	Γ	7468	4585863 NT		0.0E+00 AI244247.1					0.0E+00 BE697125.1	.00620.1	.00620.1	8489			
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00/	0.0E+00	0.0E+00	0.0E+00 B	0.0E+00	0.0E+00 B	0.0E+00	0.0E+00	0.0E+00/	0.0E+00	0.0E+00	0.0E+00 Z42399.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00E	0.0E+00	0.0E+00
	Expression Signal	8.45	8.45	1.25	1.88	1.43	. 1.43	3.49	0.94	0.94	1.1	13.6	13.6	1.98	1.02	5.35	1.29	1.29	2.47	1.21	1.06	1.43	6.29	4.71	4.71	2.79	2.79	2.04	2.04	1.7	2.76	1.41	4.46
	ORF SEQ ID NO:	12027	12028	12030	12031	12032	12033				12087	12091	12092	12093	12094	12095	12097	12098			12101		12109	12111	12112	12118	12119	12125	12126	12129			12149
	SEQ ID NO:					6933			6956	6956	6983	6987	6987	6969		6991	6993	6993	6995	6997	8669	7000	7004	7006	2006	7 1	7011	7016	7016	2019	7037	238	7040
	Probe SEQ ID NO:	1943	1943	1945	1946	1947	1947	1954	1971	1971	2000	808	2004	2006	2002	2008	2010	2010	2012	2014	2015	2017	2021	2023	2023	2028	2028	2033	2033	2036	2055	2056	2058

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Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	PM0-BT0547-210300-004-F04 BT0547 Homo sepiens cDNA	Homo septions glutarthione S-transferase theta 2 (GSTT2) and glutarhione S-transferase theta 1 (GSTT1)	genes, complete cds	IL3-CT0219-271089-022-G10 CT0219 Homo sapiens cDNA	QV-BT065-020369-092 BT085 Homo sapiens cDNA	QV-BT065-020399-092 BT065 Hamo sepiens cDNA	Human DNA-binding protein mRNA, 3'end	601122338F1 NIH_MGC_20 Homo sapiens CDINA done IMAGE: 3340068 b	Human mRNA for KIAA0244 gene, partiel cds	AV738288 CB Homo septens cDNA clone CBNBDE08 5	AV738288 CB Hamo saplens cDNA clone CBNBUEU8 5	0032e01.s1 NCI_CGAP_Lu5 Homo sapiens cLinA clone IMAGE:135/895 3	Human apolipoprotein B-100 (apoB) gene, exons 22 through 29	602014829F1 NCI_CGAP_Bm64 Homo sapiens cDNA clone IMAGE:4150734 5	601572186T1 NIH_MGC_55 Homo saplens CDNA clone IMAGE:3838012 3	CM1-TN0141-250900-439-b08 TN0141 Homo septens CDNA	CM1-TN0141-250900-439-b08 1N0141 Homo sepiens CUNA	601900261F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4128622 5	2	3540 3' similar to		5540 3' similar to			Homo sapiens chromosome 21 segment noz rover	Hamo saplens KIAA0952 protein (NIAA0952), illiniar	Homo sapiens KIAA0952 protein (KIAA0952), mKNA	Human beta-prime-edeptin (BAM22) gene, exon 16	A12b10.r1 NCI_CGAP_GCB1 Homo sapiens cunA cone IMACE:/12b1	601432317F1 NIH_MGC_72 Home sapiens curva cidne invacinity as a	Homo saplens E1A binding protein p300 (EP300) mitting	Homo sapiens KiAA0952 protein (KIAA0952), mikina	601433525F1 NIH MGC 72 Homo septens CLINA clane IMACCE 38 10007 3	
1	Oetabase Source	EST_HUMAN		M	EST HUMAN	EST_HUMAN	EST_HUMAN	TN	EST_HUMAN	NT	EST_HUMAN	EST_HUMAN	EST HUMAN	NT	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	FST HIMAN		EST HUMAN		EST_HOMAN	5	N	攴	ᅺ	TN	EST_HUMAN	EST HUMAN	TN	M	EST_HUMAN	ļ 
	Top Hit Acession No.	3E072624.1		AF240786.1	4W752708.1	41904640.1	41904640.1	14787.1	3E274696.1 E		AV738288.1 E	AV738288.1	AA931691.1	M19828.1	BF344434.1	BE748899.1	BF377897.1	BF377897.1	BF313617.1	DE018750 1	T	AA042813.1				AL163204.2	7662401 NT	7662401 NT	U36264.1	AA282281.1	BE897487.1	4557556 NT	7662401	BE895281.1	-
Most Similar	<u> </u>	0.0E+00		0.0E+00	0.0E+00	0.0E+00/	_	0.0E+00		_	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	_		0.0E+00				0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	
	Expression Signal	1.77		2.11	3.64	4.62	4.62	1.21	1.93	1.01	42.14	42.14	3.7	1.02	62.74	55.46	4.45	4.45	4.41	,	***	2.17		2.17	2.86	2.86	2.34	2.34	2.31	1.43	2.22			60.9	
	ORF SEQ ID NO:	12150		12151	12152	12154	L	1	12214		12218				12225			12230	١.	_	12238	12240		12241	12249	12250				12257					
	SEQ ID	7041		7043	7044	7046	7046	9602	7102	7104	7105	7105	7107	7109	7112	7113	7116	1		<u> </u>	77.77	7424	1	7124			L	1					L		╛
	Probe SEQ ID NO:	2059		2061	2062	2064	2064	2116	2122	2124	2125	2125	2127	2129	2132	2133	2136	2136	2140		2143	2446	C#17	2145	2153	2153	2154	2154	2150	2169	2167	2180	2185	2 6	

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Top Hit Descriptor	AU118082 HEMBA1 Homo septems cDNA clare HEMBA1002820 51	AU118082 HEMBA1 Homo serviers CDNA close HEMBA1002509 5	Homo serviens hynothetical protein Fill 1900s4 / El 19	WRO-BN0070-090800-079-8419 BN0070 House Shilliam Chair	AU19582 HEMBA1 Homo seplens cDNA close HEMBA10na155 5	000002.x1 Soares_Nit-MiPu_S1 Homo sepiens cDNA clone MAGE:1660683 3' similar to TR:008692	Homo saplens hyrothetical protein El (2002 /El (2002) - Data	601432608F1 NIH MGC 72 Homo seriens ATMA Aboo NAACE: Society F1	AB005622 HeLa CDNA (T. Noma) Homo saniens cDNA similar to adominate Management	Homo sapiens du famate receptor, ionofronio, N-methy D. esperate 24 (GRINDA)	Homo sapiens gene for cholecystokinin type A resenter complete cde	Homo sapiens gene for cholecystokinin troe-A recentor complete cris	Homo sapiens immunoalobulin-like transcript 10 variant 4 (II 11c) gans arms	602018058F1 NCI CGAP Bm67 Home seniens cDNA clone IMAGE-415970 F1	Homo sapiens collegen, type XII, alpha 1 (COL12A1), mRNA	602184558T1 NIH MGC 42 Homo septens cDNA clone IMAGE 43003R3 3"	ha04h04.x1 NCI_CGAP_Kid12 Homo sapiens cDNA clone IMAGE:2872759 3'	UHF-BP0p-els-c-07-0-Ul.rt NIH_MGC_51 Homo sapiens cDNA clone IMAGE:3072780 6	RC3-ST0197-300300-016-c04 ST0197 Homo saplens cDNA	601592530F1 NIH_MGC_7 Homo sapiens cDNA clone IMAGE:3946518 5	Hamo sapiens death receptor 6 (DR6), mRNA	UHH-BI4-acz-b-08-0-UI.s1 NCI_CGAP_Sub8 Homo saplens cDNA clone IMAGE:30865353'	Homo sapiens mRNA for membrane transport protein (XK gene)	Homo sapiens platelet-derived growth factor receptor-like (PDGFRL) mRNA	601503356F1 NIH_MGC_70 Homo septens cDNA clone IMAGE:3905148 6	Homo sapiens similar to rat Integral membrane glycoprotein POM121 (POM121L1), mRNA	RC4-HT0276-160200-013-d05 HT0276 Homo saplens cDNA	Human Sec62 (Sec62) mRNA, complete cds	601508211F1 NIH_MGC_71 Home septens cDNA clone IMAGE:3908866 5'	601489241F1 NIH_MGC_69 Homo sepiens cDNA clone IMAGE:3891371 6'	601489241F1 NIH_MGC_69 Homo sepiens cDNA clone IMAGE:3891371 5'	AF114027 Homo sapiens lung fetus Homo sapiens cDNA clone ESF6	Homo sapiens adiloan mRNA, complete ods
Top Hit Database Source	EST HUMAN	EST HUMAN	Į.	EST HUMAN	EST HUMAN	EST HUMAN	¥	EST HUMAN	EST HUMAN	IN	ΙΝ	NT	NT	EST HUMAN	NT	EST HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	EST_HUMAN	TZ	EST HUMAN			EST_HUMAN		EST_HUMAN		EST_HUMAN	EST_HUMAN	П	T_HUMAN	Ŋ
Top Hit Acession No.	AU118082.1	AU118082.1	23089	BE814424.1	AU119582.1	Al042035.1	8923620 NT	BE895605.1	Π	6006002 NT	D85606.1	D85606.1	AF106275.1	BF345274.1	29777	BF569144.1		AW501010.1	AW813853.1	BE795542.1	57038	11	232684.2	5453871 NT	3E910378.1	7657468 NT	3E150865.1	193239.1		3E875511.1		\F114027.1	F245505.1
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	-	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	-	0.0E+00	0.0E+00	-	_	0.0E+00		0.0E+00			_	0.0E+00 E	0.0E+00	1	1	0.0E+00	0.0E+00	]	151	0.0E+00
Expression Signal	21.5	21.5	1.24	1.51	0.99	3.28	1.18	4.15	3.39	5.53	1.57	1.57	1.34	0.95	6:33	16.53	2.46	2.09	2.13	27.9	1.18	1.33	3.32	3.21	1.05	2.35	0.91	3.02	2.2	4.05	4.05	1.27	=
ORF SEQ ID NO:	12391	12392	12393		12441		12442				12463	12464	12473	12477	12484	12490	12497	12498		12526	11981	12527	12529		12533	12534	12535	12536	12542	12547	12548	12549	12552
Exon SEQ ID NO:	7273	7273	7274	7291	7322	7323	7324	7328	7339	7342	7344	7344	7352	7355	7361	7369	7378	7380	7404	7409	6887	7410	7413	7415	7418	7419	7420	7421	7427	7430	7430	7431	7433
Probe SEQ ID NO:	2298	2298	2299	2316	2348	2349	2350	2354	2365	2369	2372	2372	2380	2384	2390	2398	2407	2409	2433	2438	2439	2440	2443	2445	2448	2449	2450	2451	2457	2461	2461	2462	2464

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Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	601064738F1 NIH MGC 10 Hamp september cDNA close MAR DE 3454464 F1	AU143277 Y79AA1 Homo seniens CPNA close V70A/8 40046725 E1	AU143277 Y79AA1 Homo serviews CDNA closs - V70A & 1001675 5	601105312F1 NIH MGC 15 Homo captions of NA Alexander El	601105312F1 NIH MGC 15 Homo sepiens cDNA clone IMAGE: 20870F5 FF	7q27h12x1 NCL CGAP_GC6 Homo sapiens cDNA clone IMAGE: 3' similar to TR:000246 000246 HYPOTHETICAL to 3 KD PROTEIN	Home saniens adilican mRNA American ade	601173631F1 NIH MGC 17 Hamp sabilens chivid show MAA DE 3520450 F	Homo sapiens mRNA for KIAA1415 protein partial ods	Homo saplens mRNA for KIAA1415 protein partial ods	ULH-BW1-amp-612-0-11 st NCI CGAP Sub-to-more contract of	602152653F1 NIH MGC 81 Homo seniens cDNA clare 1840. CDNA care in MGC 81 Homo seniens cDNA clare 1840.	601279873F1 NIH MGC 39 Homo sepiens cDNA close IMACE 3621786 F	Homo seplens mRNA for KIAA1321 prolem, partial cols	Homo seplens TATA box binding protein (TBP) essociated factor, RNA polymerase II, I, 28kD (TAF2I) mRNA	Homo saciens mRNA for KIAA1438 motern period cyle	601590108F1 NIH MGC 7 Home saniers CDNA charactersourser in	601590108F1 NIH MGC 7 Homo sepiens cDNA clone IMAGE: 3944304 5	601143722F1 NIH_MGC_15 Homo sapiens cDNA clone IMAGE:3051389 5	601584930F1 NIH_MGC_7 Homo saplens cDNA clone IMAGE:3939222 51	Homo sapiens mRNA for KIAA0903 protein, partial cds	Homo sapiens IMP (inosine monophosphate) dehydrogenase 1 (IMPDH1) mRNA	Homo saplens guanylate cyclase-activating protein 2 (GUCA1B) gene, exon 1	Homo sapiens mRNA for KIAA0536 protein, partial cds	AU133385 NT2RP4 Hamo sapiens cDNA clane NT2RP4001884 6"	Human bullous pemphigold antigen (BPAG1) mRNA, complete cds	AU130403 NT2RP3 Homo sapiens cDNA clone NT2RP3000779 5'	AU130403 NT2RP3 Homo sepiens cDNA clone NT2RP3000779 6	RC1-070086-220300-011-d07 OT0086 Homo sepiens cDNA	7h15h05.x1 NCI_CGAP_Co16 Homo sapiens cDNA clone IMAGE:3316089 3'	601298714F1 NIH_MGC_19 Homo saplens cDNA clone IMAGE:3628923 5'	601278373F1 NIH_MGC_39 Home saplens cDNA clone IMAGE:3610267 6'
201	Top Hit Database Source	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	LN	EST HUMAN	N <sub>T</sub>	F	EST HUMAN	EST HUMAN	EST HUMAN	LN	<del>-</del>	Ę	EST HUMAN	EST HUMAN	EST_HUMAN	EST_HUMAN	NT	TN	NT	ᅜ	EST_HUMAN	Ę	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN	EST_HUMAN
	Top Hit Acession No.	BE\$36921.1	AU143277.1	AU143277.1	BE292896.1	BE292896.1	BF223041.1	AF245505.1	BE296613.1	Γ	AB037836.1	BF513835.1			AB037742.1	5032150 NT	AB037859.1		Ī	BE293328.1		AB020710.1	4504686 NT		AB011108.1				AU130403.1	AW887015.1			3E531263.1
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00			-	0.0E+00			0.0E+00	0.0E+00	0.0E+00	-		0.0E+00	0.0E+00			+=	0.0E+00		_	_					•	0.0E+00	0.0E+00	_		0.0E+00 E
	Expression Signal	1.91	3.77	3.77	3.86	3.86	0.83	5.96	0.91	1.49	1.49	1.95	1.25	1.1	2.22	2.36	5.06	1.42	1.42	1.35	6.42	1.06	2.7	4	1.3	1.88	1.75	2.23	2.23	1.36	1.01	5.22	8.33
	ORF SEQ ID NO:	12560		12568		12570	12571	12573	12591		12609		12614		12623	12624	12626	12627	12628	12629		12637	12645	12653	12654	12657	12658	12661	12662	12665	12668	12669	
	Exon SEQ ID NO:			7453	7454	7454	7455	7458	7476	7708	7708	7489	7494	7496	7504	7505	7507	7508	7508	7511	7518	7520	7527	7537	7540	7543	7544	7547	7547	7550	7553	7554	7555
	Probe SEQ ID NO:	2480	2485	2485	2486	2486	2487	2490	2508	2522	2522	2523	2528	2530	2539	2540	2542	2543	2543	2546	2553	2555	2563	2574	2577	2580	2581	2584	2584	2587	2290	2591	2592

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Homo seplens hypothetical protein F. 111052 /F. 111052) m.D.N.A.	Homo seciens mRNA for KIAA1311 protein perital cass	EST188414 HCC cell line (matestasts to liver in mouse) II Homo sepiens cDNA 5' end skniker to ribosomal profein   20	601589625F1 NH MGC 7 Homo sentens cDNA clara MACE: 3043E04 E1	Human beta-brime adendin (BAM22) gene evan 6	Homo saplens neurogulin 1 (NRG1) transcript varient SMDF mRNA	Homo sepiens skeletal muscle LIM-protein 1 (FHI 1) gene complete ode	Homo sapiens hG28K mRNA for GTP-binding protein like 1 complete cole	601591991F1 NIH MGC 7 Homo sapiens cDNA clone IMAGE:3945983 5	602155923F1 NIH MGC 83 Homo sapiens cDNA clone IMAGE-4297132 F	601335485F1 NIH MGC 39 Homo sapiens CDNA clane IMAGE:3689584 F	AV721647 HTB Homo sapiens cDNA clone HTBBYE09 5'	Homo sapiens spermatogenests associated PD1 (KIAA07571 m.PNA	Homo sapiens spermatopenesis associated PD1 (KIAA0757) mBNA	Homo saplens hypothetical protein FL 120477 (FI 120477) mRNA	Homo saplens hypothetical protein FLJ20477 (FLJ20477) mRNA	Homo sapiens hypertension-related calcium-regulated ones mRNA complete calc	AV651066 GLC Homo sepiens cDNA clone GLCC D07 3'	CM1-TN0141-250900-439-b08 TN0141 Homo septems cDNA	CM1-TN0141-250900-439-b08 TN0141 Homo saplens cDNA	601560903F1 NIH_MGC 9 Homo sapiens cDNA clone IMAGE:3829472 5	601462038F1 NIH_MGC_68 Hamo sepiens cDNA clane IMAGE:3865497 6		mo sapiens cDNA clone IMAGE:3071340 3'				602085579F1 NIH_MGC_83 Hamo sapiens cDNA clane IMAGE:4246918 5	Homo sapiens protein tyrosine phosphatase, receptor type, T (PTPRT), mRNA			IA clane IMAGE:2518663 5' similar to
 Top Hit Database Source	M	N	EST HUMAN	EST HUMAN	NT.	NT	N	Z.	EST_HUMAN	EST_HUMAN	EST HUMAN	Г						EST HUMAN	Т	Т	Г	EST_HUMAN	ĽΝ	EST_HUMAN				EST_HUMAN			EST_HUMAN	EST_HUMAN
Top Hit Acession No.	8922843 NT	AB037732.1	AA316723.1		U36253.1	669517	AF110763.1	AB051826.1	BE796376.1	BF680632.1	BE563433.1	AV721647.1	5174486 NT	5174486 NT	8923441 NT	8923441 NT	4F290195.1	4V651066.1		3F377897.1	3E747193.1	3F037713.1	4L163201.2	3F514110.1	4503098 NT	7705275 NT	TN05275	3F677694.1	27522		V725534.1	0.0E+00 AIB79183.1
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
Expression Signal	1.38	1.22	27.94	1.31	5.13	5.03	2.44	1.27	26.29	1.5	24.53	2.28	2.82	2.82	1.27	1.27	2.26	51.51	3.33	3.33	19.56	3.09	0.93	2.53	2.14	1.05	1.05	2.68	1.43	26.73	26.73	11.75
ORF SEQ ID NO:	12690	12698		12721	12726				12735	12736	12740		12743	12744	12745	12746	12747		12748	12749	12756	12760		12768		12776	12777	12778	12786	12789	12790	
SEQ ID NO:		7586	7610	Ш		7616			7623	7624	8777	7627	7629	7629	7630	7630	7631	7632	7633	7633	7640	7645	7653	7654	7660	7665	7665	9992	7672	7676	7676	7678
Probe SEQ ID NO:	2616	2626	2650	2651	2655	2656	2657	2658	2664	2665	2668	2669	2671	2671	2672	2672	2673	2674	2675	2675	2682	2687	2695	2696	2703	2708	2708	2709	2715	2719	2719	2721

Page 193 of 209 Table 4 Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	602071957F1 NCI_CCAP_Bm67 Homo sepiens cDNA clone IMACE: 4214679 6	601450912F1 NIH MGC 65 Homo sapiens cDNA done IMAGE:3854842 5	AU131494 NT2RP3 Homo septens CDNA clone NT2RP3002672 5	Allisaddod NT 28D3 Homo sanians cDNA clone NT 2RP3002672 5	ACISTACT NICE AND ARC 17 Home series CDNA clare IMAGE:2960806 5	CONSTANT SET INITIATION OF THE SECOND CONTRACT TO THE WAGE TO THE SECOND SET INITIATION OF THE SECOND SET INITIATION OF THE SECOND SET INITIATION OF THE SECOND SET INITIATION OF THE SECOND SET INITIATION OF THE SECOND SET INITIATION OF THE SECOND SET INITIATION OF THE SECOND SET INITIATION OF THE SECOND SET INITIATION OF THE SECOND SET INITIATION OF THE SECOND SET INITIATION OF THE SECOND SET INITIATION OF THE SECOND SET INITIATION OF THE SECOND SET INITIATION OF THE SECOND SECOND SET INITIATION OF THE SECOND SECON	600944/94TI NIT MICE IT FIGURE SQUARES COLD CONTROL OF SURES IT	glycoprotein D-Duny group anugen (numer, mod, commission of commission of complete cds	Homo sapiens B I RCP2 many in 1-box and Worldware process to the process of the p	Homo sapiens ALK-ike protein mitting, per usa cus	Homo sapiens ALK-tike procein mixtva, parival cus	Homo sapiens cytochrome F400, subtamily I (atourning buypepede 1 (atourning), polypepede 1 (atourning) (CYP1B1) mRNA	Homo sapiens cytochrome P450, subfamity I (dioxin-Inducible), polypeptide 1 (glaucoma 3, primary infantile)	(CYP1B1) mRNA	H. sapiens serine hydraxymethyftransferase pseudogene	Homo sapiens 5-aminolevulinate synthase 2 (ALAS2) gene, complete cas	Homo sapiens mRNA for KIAA1527 protein, partial cds	Lorent Annie merital mil name for ribosomal protein L3, U82 snoRNA, U83a snoRNA and U83b snoRNA	Portion additions that the light of the contract of the contra	Hamo saplens chramosome 21 segment HSZ1CWU1	Human sodium channel mitNA	Human AHNAK nucleoprotein mirnA, 9 ena	H. saplens id3 gene for HLM type uanscriptum i rever	Home sepiens chromosome 21 segiment not 10000	Homo saperts zinc miger protein ZZ I (ZZ I ZZ I), milk zinc	Homo septens zinc imger protein 22 (CM 22 ), III CM	Homo sapiens zirci inga pionali za (za za za za za za za za za za za za za z	Human transglutaminase mixiva, compress cus	Homo saplens gammma-cymplasmic acum (Action 9) posucogum	Homo saplens gammma-cytoplasmic acum (ACT of 5) pecuadores	Nove human mRNA from chromosome 1, withclines entities to contact the contact to contact the contact to contact the contact to contact the contact to contact the contact to contact the contact to contact the contact to contact the contact to contact the contact to contact the contact to contact the contact to contact the contact to contact the contact	H. sepiens mRNA for nuclear DNA helicase II	Homo sapiens protocadherin alpha C1 (PCLDH-alpha-C1) innvity, white con-	Homo saplens eukaryotic translation etongation racinal	
	Top Hit Database Source	EST HUMAN	EST HIMAN	ENT HIMAN	1000	ESI HUMAN	EST HUMAN	EST HUMAN	NT	N <sub>1</sub>	NT.	17	— <b> </b>		5	ĽŽ	LZ	FZ		N.	NT	NT	NT	Z	٦	Ę	Z.	Į.	N	۲	NT	NT	12	LN	¥	
	Top Hit Acesslon No.	530661.1	070768 4	1	T	1		300344.1				0.0E+00 AF264750.1	4503202 NT		4503202 NT	0.0E+00 X85980.1		0.00±00 App.000 4	Ī	0.0E+00 AJZ38852.1		0.0E+00 M91803.1	V80902.1	C73428.1	AL16326			7019584 NT	M98478.1	D50657.1	D50657.1	AL096857.1		AF15230	4503470 NT	
	Most Similar (Top) Hit BLAST E Value	0 0E+00 BF	200	0.00	0.0=+00	0.0E+00/	0.0E+00	0.0E+00	0.0E+00 S7	0.0E+00	0.0E+00	0.0E+00	00+300	20.70	0.0E+00	0.0E+00	00130	0.05	0.0=+00				0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00		0.0E+00		1	ļ		L	
-	Expression Signal	277	189	3 3	2.2	2.2	68.47	68.47	2.82	3.43	1.92	1.92	97.6	2.10	2.78	5.17	10,	\$ 1	1.35	1.16	235			1.42	2.78	1.41	1.41	1.4.1		,				1.14		
	ORF SEQ ID NO:	42705	15/30	12/96	12797	12798	12799	12800			10768		50,7	1	11063		1				12813					12821			L					1		12634
	Exon SEQ ID NO:	7004			7684	7684	7685	7685	١.			L		6032	6043				7786	7792					1	1					1				1	8 7818
	Probe SEQ ID NO:	2000	47/7	2725	2727	2727	2728	2728	2734	2737	2743	2743		2747	7776	1417	70/7	2763	2765	1.66	2777	2775		2781		2784	278	3,000	278	2707		7617	CR/2	2707	Ž,	2798

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo seplens eukaryctic translation elongation factor 1 alpha 1 (EEF1A1) mRNA	Homo sepiens seminarthrecarine kinese 9 (STK9) mRNA	DKFZp586C0621 rf 586 (syranym; hute1) Hamp sapiens cDNA clane DKFZp58eC0624	Homo sepiens chandrolitin suifate proteccifican 4 (melanome essociated) (CSPG4), mRNA	QV2-BT0636-130400-138-h03 BT0636 Homo expiens cDNA	QV2-BT0636-130400-138-h03 BT0636 Homo sepiens cDNA	Homo sepiens low density lipoprotein-related protein 2 (LRP2), mRNA	Homo sapiens low density lipoprotein-related protein 2 (LRP2), mRNA	Homo sapiens chromosome 21 segment HS21C006	Homo sapiens chromosome 21 segment HS21C006	z96b11.s1 NCI_CGAP_GCB1 Homo sepiens cDNA clone IMAGE:683517 3' similar to contains Alu	repetuve etement. Homo earlane bHh5 seme for helr basetin associated to 0	TOTAL SOURCE IN THE TOTAL MANUAL COURS IN U.S.	Homo sapiens EpiA4 (EPHA4) mRNA	Homo sapiens eukaryotic translation elongation factor 1 alpha 1 (EEF1A1) mRNA	br18d07.x1 NCI_CGAP_Bm25 Homo sepiens cDNA done IMAGE:2167981 3' similar to TR:016247 016247 F44E7.2 PROTEIN.:	b18407.x1 NCJ_CGAP_Bm25 Homo saptens cDNA clone IMAGE:2167981 3' similar to TR:016247 016347 F44E7.2 PROTEIN.:	ZINC FINGER PROTEIN 132	Homo sapiens protocadherin gamma C4 (PCDH-gamma-C4) mRNA, complete cds	Homo sapiens mRNA for KIAA1267 protein, partial cds	Homo sapiens mRNA for KIAA1267 protein, partial cds	Homo sapiens mRNA for KIAA1508 protein, partial cds	Homo sapiens mRNA for KIAA1508 protein, partial cds	Homo sapiens KIAA0100 gene product (KIAA0100), mRNA	Homo sepiens KIAA0100 gene product (KIAA0100), mRNA	Homo septens myeloid/lymphoid or mixed-lineage feukennia (trithorax (Drosophila) homolog); translocated to, 4 (MLLT4) mRNA	Homo sepiens myeloid/fymphold or mised-lineage leukemia (trithorax (Drosophila) homolog); transtocated to, 4 (MLLT4) mRNA	7n40403.x1 NCI_CGAP_Lu24 Homo sepiens cDNA clone IMAGE:3567028 3' similar to TR:QBVLN1 Q9VLN1 CG17293 PROTEIN.;	
	Top Hit Database Source	5	Į.	EST HUMAN		T HUMAN	EST_HUMAN		둗	TZ.	N-1		ESI HOMAN			F.	EST HUMAN		Т	Г		NT	FX	NT		NT	:		EST_HUMAN	
	Top Hit Acession No.	4503470 NT	4507280 NT	AL047599.1	4503098 NT	BE081896.1	BE081896.1	6806918 NT	5806918 NT	AL163206.2	AL163206.2	A A A 4 E E 7 O 4			4758279NI	4503470 NT	Al561002.1		P52740	AF152338.1		AB033093.1	AB040941.1	AB040941.1	7661903 NT	7681903 NT	6174574 NT	5174574 NT	BF110702.1	
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	200	_		0.0=+00	0.0E+00	0.0E+00			0.0E+00		0.0E+00			0.0E+00	0.0E+00	0.05+00	0.0E+00	0.0E+00 E	
[	Expression Signal	71.64	2.53	-	2.12	5.88	5.88	0.85	0.85	3.1	3.1	4 03	4.09	2	30.0	50.46	2.63	2.63	1.19	1.05	2.2	2.2	5.42	5.45	2.79	2.79	3.2	3.2	0.99	
	ORF SEQ ID NO:		12846	12850		12852			12862	12866	12867	42868		42075	12070	128/6	12877	12878	12880	12881	12893	12894	12895	12896	12899	12900	12901	12902	12906	
	Exan SEQ ID NO:	7818	7830	7833	7834	7837	7837	7843	7843	7846	7846	7847	7854	7958	7967	3	7858	7858	7860	7861	7877	7877	7878	7878	7881	7881	7882	7882	7886	
	Probe SEQ ID NO:	2798	2810	2813	2814	2817	2817	2822	2822	2825	2825	2826	2834	2826	2007	/287	2838	2838	2840	2841	2857	2857	2858	2858	2861	2861	2862	2862	2867	

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	7n40d03.x1 NCI_CGAP_Lu24 Hamo sapiens cDNA done IMAGE:3567026 3' similar to TR:Q9VLN1 Q9VLN1 CG17263 PROTEIN:	Homo sapiens melanoma antigen, family B, 4 (MAGEB4), mRNA	Homo sepiens melanoma antigen, family B, 4 (MAGEB4), mRNA	Homo sapiens neurewin III (NRXN3) mRNA	H.sapiens mRNA for M phese phosphoprotein 10	Homo sapiens mRNA for KIAA1208 protein, partial cds	H.sapiens NF-H gene, exon 4	H.sapiens NFH gene, exon 4	Homo saplens immunoglobulin-like transcript 1c variant 4 (ILT1c) gene, exon 6	qf43f09.xf Soares_lestis_NHT Homo appiens cDNA clone IMAGE:1752809 3"	Homo sapiens prospero-related homeobox 1 (PROX1) mRNA	Homo sapiens mRNA for PKU-elpha, partial ods	Hamo sapiens KIAA0737 gene product (KIAA0737), mRNA	Homo sapiens calcium channel, voltage-dependent, gamma subunit 3 (CACNG3), mRNA	Homo sapiens calcium channel, voltage-dependent, gamma subunit 3 (CACNG3), mRNA	Homo sapiens intersectin short isoform (ITSN) mRNA, complete cds	Homo sapiens intersectin short isoform (ITSN) mRNA, complete cds	Human displacement protein (CCAAT) mRNA	EST388375 MAGE resequences, MAGN Homo sepiens cDNA	Homo sapiens membrane-bound aminopeptidase P (XNPEP2) gene, complete cds	Homo sapiens heat shock 70kD protein 1 (HSPA1A), mRNA	Homo sapiens heat shock 70kD protein 1 (HSPA1A), mRNA	soform 2 of a novel human mRNA from chromosome 22	Homo sapiens putative transcription factor CR53 (CR53) mRNA, partial cds	Homo sepiens transcription factor (GHM enhancer 3, JM11 protein, JM4 protein, JM5 protein, T54 protein,	JM10 protein, A4 differentiation-dependent protein, triple LIM domain protein 6, and synaptophysin genes,	complete cds; and L-type calcium chemol a>	Human gemiline gene 16.1 for lg lembda L-chain C region (IgL-C18.1)	Homo sapiens F-box protein FBL5 (FBL5) mRNA, complete cds	Homo sapiens melanoma-associated antigen (MAGE-C1) gene, complete cds	Homo sepiens SW I-SNF complex protein p270 mRNA, pertial cds	Homo saplens NOD1 protein (NOD1) gene, exons 1, 2, and 3	Homo sepiens KIAA0469 gene product (KIAA0469), mRNA
Top Hit Database Source	EST HUMAN	N	¥	¥	TN	M	N	TN.	N	EST_HUMAN	5	Z	5	5	レフ	4	N	TN LN	EST_HUMAN	VT.	F	느	NT	NT			NT	NT	Ρ	NT	NT	NT	トフ
Top Hit Acession No.	3F110702.1	4505084 NT	4505084 NT	4758827 NT	98494.1	0.0E+00 AB033034.1		(15309.1	_		4506118 NT	B004884.1	7662273 NT	5729755 NT	5729755 NT	F114488.1	F114488.1		L	F195953.1	5579469 NT	5579469 NT	L359403.1	F017433.1			F196779.1	03529.1	F199355.1		F265208.1	F149773.1	7682139 NT
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 X15309.1	0.0E+00)	0.0E+00	0.0E+00 AI149880.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00 M74099.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00			0.0E+00	0.0E+00	0.0E+00 A	0.0E+00	0.0E+00	0.0E+00	0.0E+00
Expression Signal	0.99	2.17	2.17	1.65	1.14	0.93	0.94	0.94	8.37	1.38	1.21	4.29	2.03	2.05	2.05	1.3	1.3	1.23	0.84	4.06	6.1	6.1	6.5	1.96		,	1.97	3.21	1.61	1.43	3.49	5.9	3.23
ORF SEQ ID NO:	12907	12917	12918	12923		12926	12927	12928	12930		12950	12951	12959	12960	12961	12971	12972	12992	12999		13005	13006		13010				13033		13040	13059	13060	13063
Exon SEQ ID NO:				7901	7902		7907		7909			7832			ı		7954	7977	7985	7988	7991	7891	7993	7997	-		8	8020	8025	8029	8050	8051	8055
Probe SEQ ID NO:	2867	2875	2875	2882	2883	2886	2888	2888	2890	2904	2912	2913	2924	2925	2925	2935	2935	2958	2967	2970	2973	2973	2975	2979			2982	3002	3008	3012	3033	3034	3038

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Table 4
Single Exon Probes Expressed in HBL100 Cells

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	Top Hit Descriptor	Homo saplens offactory receptor-like protein (OLFR 42B) cense. Of FR 42B and 10 allake madial and	form saniens noticeshim unhang notice of the salas and s	Hunan fertiin heavy chain mRNA complete calc	Homo sapiens mRNA for KIAA0549 prohein partial cys	Homo sapiens mRNA for KIAA0549 protein, partial cds	ye3203 s1 Stratagene lung (#837210) Homo sapiens dDNA dane IMAGE:119453 3' similar to SP:S29539 S29539 BASIC PROTEIN 23K .	601878507F1 NIH MGC 55 Homo sapiens cDNA clove IMAGE 4107433 F	wu12h10x1 NCI CGAP GCB Homo seniens cDNA claws MAAGE 3568073 31	H. sapiens mRNA for gemme-glutamytransferase	H.sapiens mRNA for gamma-glutamytransferase	Homo sapiens neureach III (NRXN3) mRNA	Homo sapiens neurexin III (NRXN3) mRNA	Homo sapiens interleukin 1 receptor, type I (IL1R1) mRNA	Homo sepiens nucleolar phosphoprotein B23 (NPM1) mRNA, complete cds	Homo sapiens solute carrier family 25 (mitochondrial carrier, adentire nucleotide translocator), member 6 (SLC2SAS), nuclear gene encoding mitochondrial protein mRNA	Homo saplens CREB binding protein (Rubinstein-Taybi syndrome) (CRERAD) mRNA	Homo sepiens CREB binding protein (Rubinstein-Taybi syndrome) (CREBBP) mRNA	ae87b11.s1 Stratagene schizo brain S11 Homo sapiens cDNA clone IMAGE:971133 3'			Homo sapiens fibrillin 1 (Marfan syndrome) (FBN1) mRNA		Homo sapiens HLA class III region containing tenescin X (tenescin-X) gene, partial cds; cytochrome P450 21-	hydroxylase (CYP21B), complement component C4 (C4B) G11, helicese (SKI2W), RD, complement factor B	(b), and comprehensial comparison of (cc) games,	(VLGR1) mRNA, complete cds		Homo sapiens KIAA0440 protein (KIAA0440), mRNA	Homo sapiens A kinase (PRKA) anchor protein 1 (AKAP1), mRNA		Homo sapiens SWI-SNF complex protein p270 mRNA, partial cds
20001	Top Hit Database Source	NT	- 5	Į.	Z.	TN	EST HUMAN	EST HUMAN	EST HUMAN	TN	LN	TN.	7	Ę	N	<b>-</b>	7	7	EST_HUMAN	TN	LN L	トア	<b>ال</b>		<u>-</u> <u>-</u> -		LN		Ę	1		I.V.
	Top Hit Acession No.	AF042075.1	4826783 NT	120941.1	-:	AB011121.1	T94870.1	BF243336.1	Al968086.1	X98922.1	X98922.1	4758827	4758827 NT	4504658 NT	M28699.1	4502098 NT	4758055 NT	4758055 NT	AA774783.1	AF286598.1	AF286598.1	4557590 NT	4507720 NT	-	AE040449 4	Ī	AF-055084.1	7662125 NT	7662125 NIT	4502014 NIT		4F265208.1
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00		00110		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
	Expression Signal	1.38	3.44	28.05	1.09	1.09	33.16	1.16	1.26	4.08	4.08	1.48	1.48	9.09	12.76	10.04	8.0	9.0	2.91	5.5	5.5	1.67	0.98		1 87	5	3.59	1.14	1.14	2.89	2.89	2.91
	ORF SEQ ID NO:	13064	13098	13107		13111	13118		13137		13145				13182	13185	13193	13194	13195	13203	13204	13212	13219		13225	220	13228	13230	13231	13238	13239	13254
	Exon SEQ ID NO:	8056	8084	8093	8096	8096	8103	8118	8120	8125	8125	8136	8136	8143	8161	8165	8171	8171	8173	8181	8181	8191	8196		8203		8290	8078	8208	10045	10045	8233
	Probe SEQ ID NO:	3039	3068	3077	3080	3080	3087	3102	3104	3109	3109	3120	3120	3127	3145	3149	3155	3155	3157	3165	3165	3175	3180		3187		0815	3183	318	3201	3201	3218

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Single Exon Probes Expressed in HBL100 Cells

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Top Hit Descriptor	Homo seplens hybothetical protein FI. (20895/FFI. (20805), mBNA	Homo sepiens douth receptor 6 (DR6), mRNA	#58f08.22 NG_CGAP_Part Homo explens cDNA done IMAGE:2222555 3' similar to SW:RL11_RAT P25121 60S RIBOSOMAI PROTEIN 141	Homo sapiens relamenese neverse transcrimene (TERT) come arrows 1.8	Homo saniens felomenese reverse transcriptose (TEDT) seves account 4 a	Homo sapiens hormonally upresulated neu hymoressociated kinese (HI NKY) mBN/s	Homo Saciens hormonally uprequiated neu humor-essociated kinase (HI NIC). The American	Homo saplens caspase 8, apoptosis-related cysteine proteese (CASPR) mRNA	Homo seplens caspase 8, apoptosis-related cysteine professe (CASPR) mRNA	Homo saplens pyrin (MEFV) gene, complete cds	Homo sapiens mRNA for KIAA 1507 protein partial cols	wb10104.xt NCI_CGAP_GC6 Homo septiens aDNA alone IMAGE:2305279 3' similar to TR:Q81929 Q61929 ZINC FINGER PROTEIN	AU123664 NT2RM2 Homo saplens cDNA clone NT2RM2000735 57	Homo sapiens offactory receptor, femily 10 subfamily C member 1 (OB40C4) mBNA	Homo sapiens offactory receptor, family 10, subfamily C member 1 (OR10C1), mRNA	Homo saplens neuroblastoma-amplified protein (LOC51594), mRNA	Homo saplens T-twoe calcium channel airthart schaint Afrikaria isoform (CACNA1) mDNA	MR1-SN0033-100400-001-008 SN0033 Home sexiens CDNA	Homo sapiens KIAA0952 protein (KIAA0952), mRNA	Homo sapiens KIAA0952 protein (KIAA0852), mRNA	Homo saplens beaded filement structural protein 1, filensin (BFSP1) mRNA	Homo sepiens leukocyte immunoglobulin-like receptor, subfamily A (with TM domain), member 2 (LILRA2), mRNA	Homo sapiens skeletal muscle LIM-protein 1 (FHL1) gene, complete cds	Homo sapiens death receptor 6 (DR6), mRNA	Homo sapiens protein kinase, AMP-activated, eipha 2 catalytic subunit (PRKAA2) mRNA	Homo sapiens protein kinase, AMP-activated, alpha 2 catalytic subunit (PRKAA2) mRNA	Homo sapiens mRNA for rape-2 (rape gene)	Homo sapiens mRNA for rapa-2 (rapa gene)	Bectertophage P1 replication region including repA, parA, and parB genes and inoA, inoB, and incC incompatibility determinants
Top Hit Database Source	¥	Z.	EST HUMAN	Т	LN L	¥	¥	Ę		Ę		EST HUMAN	Т	1			Ż	T_HUMAN					F.					TN	I N
Top Hit Acession No.	8923624 NT	7657038 NT	0.0E+00 AI589294.1		T	7657213	7657213 NT	4502582 NT	4502582 NT	0.0E+00 AF111163.1	Г		J123664.1	7363436	7363436 NT	7706239 NT	0.0E+00 AF211189.1		2401	7662401 NT	4502398 NIT	5803067 NT	F110763.1	7657038 NT	5453965 NT	5453965 NT	1277276.1	1277276.1	
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00 A	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 AI	0.0E+00	0.0E+00	0.0E+00	0.0E+00[A.	0.0E+00 A.	0.0E+00 K02380.1
Expression Signel	1.97	1.1	9	2.09	2.09	0.78	0.78	1.16	1.16	11.21	1.25	0.86	3.08	0.95	0.95	1.91	1.03	76.0	1.39	1.39	1.13	2.13	1.45	2.22	1.37	1.37	96.0	96.0	4.78
ORF SEQ ID NO:		13268	13284	13292	13293	13294		·	13298	13300	13302	13316	13346	13349	13350	13352	13353		13366	13367	13368	13369	12728	13382	13383	13384	13388	13389	13390
Exen SEQ ID NO:	8234	8246	8263	8270	8270	8271	8271	8274	8274	8277	8279	8291	8323	8329	8329	8332	8333	8337	8349	8349	8350	8352	7817	8365	8366	8366	8369	8369	8370
Probe SEQ ID NO:	3219	3231	3250	3257	3257	3258	3258	3261	3261	3264	3266	3279	3312	3319	3319	3322	3323	3327	3340	3340	3341	3343	3352	3357	3358	3358	3361	3361	3362

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Single Exon Probes Expressed in HBL100 Cells

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Single Exon Probes Expressed in HBL100

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Top Hit Descriptor		Homo sapiens sel (Drosophila)-Rice 1 (SALL1), mRNA	ox77c11x1 Soares_NhtHMPu_S1 Hamo septens cDNA clone IMAGE:1882356 3' similar to WP:T1984.4 CE13742;	Homo sepiens butyrophilin, subfamily 3, member A3 (BTN3A3), mRNA	QV0-CT0225-230300-169-e01 CT0225 Homo seplens cDNA	Homo sepiens H3 histone family, member K (H3FK), mRNA	Homo sapiens gamme-glutamykcysteine synthetase (GLCLC) gene, partial cds	602084563F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4248598 5	oq94h06.s1 NCI_CGAP_Kid6 Homo septens cDNA cione IMAGE:1594043 3' similar to contains MER29.b2 MER29 repositive element	QV0-DT0047-170200-123-g01 DT0047 Homo sepiens cDNA	602152486F1 NIH MGC B1 Homo saplens cDNA dome IMAGE-42038455	602152486F1 NIH MGC 81 Hamo saciens cDNA clone IMAGE:4293645 5	Homo saciens retinoblastoma-binding protein 2 (RBBP2) mRNA	184g01.x1 Soares, NFL, T, GBC, S1 Homo sapiens cDNA clone IMAGE: 297g024.31	hi84g01.x1 Sogres_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2979024.3'	Homo sepiens KIAA0808 gene product (KIAA0808), mRNA	Homo sepiens midline 1 (Opitz/BBB syndrome) (MID1) mRNA	Homo sapiens midline 1 (Opitz/BBB syndrome) (MID1) mRNA	Homo sapiens mRNA for G protein-coupled inward rectifier potassium channel, complete cds	Homo sapiens glyceraldehyde-3-phosphate dehydrogenase (GAPD), mRNA	Homo sapiens WAVE2 mRNA for WASP-family protein, complete cds	Homo sapiens mRNA for KIAA0406 protein, partial cds	Homo sapiens SH2-containing protein Nsp2 mRNA, complete cds	Homo sapiens SH2-containing protein Nsp2 mRNA, complete ods	NHTBCae15g09f1 Normal Human Trabecular Bone Cells Homo sapiens cDNA clane NHTBCae16g09	NHTBCee15g09f1 Normal Human Trabscular Bone Cells Homo sapiens cDNA clone NHTBCee15g09	Homo sapiens chromosome 21 segment HS21C004	Homo sapiens chromosome 21 segment HS21C004	Human mRNA for KIAA0333 gene, partial cds	Human mRNA for KIAA0333 gene, partial cds	MR2-CT0222-281089-005-605 CT0222 Homo sapiens cDNA
Top Hit Database Source	- 1	NT	EST HUMAN	N.	EST_HUMAN	LN FN	N L	EST_HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	EST HUMAN	N.	EST HUMAN	EST HUMAN	NT.	FX	Ę	LN	7	NT	NT	TN	TN	EST_HUMAN	EST HUMAN	LN	NT	N	NT	EST_HUMAN
Top Hit Acessian No.		6997248 NT	A1081907.1	6325463 NT	0.0E+00 AW852217.1	4504294 NT		0.0E+00 BF676383.1	0.0E+00 AA988715.1	0.0E+00 AW937977.1	0.0E+00 BF672054.1		4826967 NT	0.0E+00 AW664693.1	0.0E+00 AW664693.1	7662319 NT	4557752 NT	4557752 NT		7669491 NT			0.0E+00 AF124250.1		0.0E+00 AA852743.1	0.0E+00 AA852743.1	0.0E+00 AL163204.2				0.0E+00 AW851714.1
Most Similar (Top) Hit BLAST E	vaiue	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00/	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 D87327.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
Expression Signal		0.94	1.7	1.01	4.62	1.42	5.83	7.79	1.01	1.4	96.0	96.0	1.13	0.75	0.75	0.72	2.09	2.09	2.29	14.47	3.35	1.92	. 4.2	4.2	1.04	1.04	1.83	1.83	0.93	0.93	1.43
ORF SEQ ID NO:		13516		13519		13527		13528		13539	13547	13548	-	13550	13551	13555	13560	13561	13574		13594	13595	13596	13597	13605	13606	13608	13609	13610	13611	13614
Exon SEQ ID NO:		8501	8502					8517	8521	8529	8541	8541	8542	8544	8544	8547	8553	8553	8569	8573	8290	8591	8593	8593	8599	8599	8602	8602	8603	8603	8608
Probe SEQ ID NO:		3493	3494	3496	3500	3504	3508	3509	3513	3522	3535	3535	3536	3538	3538	3541	3546	3546	3562	3566	3583	3584	3586	3586	3592	3592	3595	3595	3296	3596	3599

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Top Hit Descriptor	Homo sepiens metalt metalloproteinase 24 (membrane-inserted) (MMP24), mRNA	rotein, partial cds	TRANSCRIPTION REGULATOR PROTEIN BACH1 (BTB AND CNC HOMOLOG 1) (HA2303)	rotein, partial cds	otein, pertial cds	ase (ASK) mRNA	ASS (ASK) MRNA	UI-H-BW0-ajs-6-12-0-UI.s1 NCI_CGAP_Sub6 Homo sapiens cDNA clone IMAGE:2733022 3'	UI-H-BW0-ajs-6-12-0-UI.s1 NCI_CGAP_Sub6 Homo septiens cDNA clone IMAGE:2733022 3'	chain, exen 6	8806g01.r1 Soares, NhHMPu, S1 Hamo sepiens cDNA clone IMAGE:812496 67 similar to SW:KRB4, SHEEP P02445 KERATIN, HIGH-SULFUR MATRIX PROTEIN, IIIB4, [1]:	otein, partial ods	Homo sapiens similar to rat integral membrane glycoprotein POM121 (POM121L1), mRNA	t (KIAA0569), mRNA	PS2) mRNA	Homo sapiens v-ets avian erythroblastosis virus E26 oncogene related (ERG), mRNA	Homo sapiens wets awan erythroblastosis wrus E26 oncogene related (ERG), mRNA	rotein (MLH3) gene, complete ods	R208) gene, partial cds	Homo sapiens similar to rat Integral membrane glycoprotein POM121 (POM121L1), mRNA	Homo sapiens similar to rat integral membrane glycoprotein POM121 (POM121L1), mRNA	Homo sapiens smooth muscle myosin heavy chain SM1 mRNA, alternatively spliced, partial cds	noogene family (RAB9) mRNA	71) gene, partial cds	71) gene, partial ods	te62f10.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clane IMAGE:209130731	CDH-beta3) mRNA, complete ods	I) (DSP) mRNA	Homo sepiens ATP-sensitive inwerdly rectifying K-channel subunit (KCNJ&BIR1) gene, complete cds	xein 2 (MECP2), mRNA	rt (KIAA0569), mRNA	Homo sapiens myosin light chain kinase Isoform 2 (MLCK) mRNA, complete cds
•	Homo sapiens matrix metalloproteinas	Homo sapiens mRNA for KIAA0796 protein, partial cds	TRANSCRIPTION REGULATOR PR	Homo sapiens mRNA for KIAA0910 protein, partial cds	Homo sapiens mRNA for KIAA0910 protein, pertial cds	Homo saplens activator of S phase kinase (ASK) mRNA	Homo saplens activator of S phase kinase (ASK) mRNA	UI-H-BWO-ajs-0-12-0-UI.s1 NCI CGA	UI-H-BW0-ajs-0-12-0-UI.s1 NCI_CGA	Human gene for Type XIX collagen a1 chain, exon 6	880801.1 Soares NhHMPu_S1 Hon SW:KRB4 SHEEP P02445 KERATIN	Homo sapiens mRNA for KIAA0903 protein, partial cds	Homo sapiens similar to rat integral me	Homo sapiens KIAA0569 gene product (KIAA0569), mRNA	Homo sapiens ribosomal protein S2 (RPS2) mRNA	Homo sapiens vets avian erythroblaste	Homo sapiens v-ets avian erythroblasts	Homo sapiens DNA mismatch repair protein (MLH3) gene, complete cds	Pan troglodytes offactory receptor (PTR208) gene, partial cds	Homo sapiens similar to rat integral me	Homo sapiens similar to rat integral me	Homo sapiens smooth muscle myosin	Homo sapiens RAB9, member RAS oncogene family (RAB9) mRNA	Gorilla gorilla offactory receptor (GGO71) gene, partial ods	Gorilla gorilla offactory receptor (GGO71) gene, partial ods	te62f10.x1 Soares_NFL_T_GBC_S1 I	Homo sapiens protocadherin beta 3 (PCDH-beta3) mRNA, complete ods	Homo septens desmoplakin (DPI, DPII) (DSP) mRNA	Homo sepiens ATP-sensitive inwerdly	Homo sapiens methyl CpG binding protein 2 (MECP2), mRNA	Homo sepiens KIAA0569 gene product (KIAA0569), mRNA	Homo sapiens myosin light chain kinas
Top Hit Database Source	Ę	F	SWISSPROT	Z	۲	Ę	NT	EST_HUMAN	EST_HUMAN	TN	EST HUMAN	Į.	Ā	TN	NT	INT	NT	ĮN	NT	IN	IN	N	NT	NT	NT	EST_HUMAN	IN	IN	토	LN	NT	Ä
Top Hit Acession No.	5729928 NT	B018339.1			0.0E+00 AB020717.1	5729733 NT	5729733 NT	W298134.1	W298134.1		0.0E+00 AA463859.1	0.0E+00 AB020710.1	7657468 NT	7662183 NT	4506718 NT	1N 5901391	7657065 NT	F195658.1	0.0E+00 AF179733.1	7657468 NT	7657468	F020091.1	4759011 NT	0.0E+00 AF127851.1	0.0E+00 AF127851.1	0.0E+00 Al377699.1	F152498.1	4758199 NT	0.0E+00 S78685.1	TN10148 NT	7662183	F069601.2
Most Similar (Top) Hit BLAST E Valtie	0.0E+00	0.0E+00 A	0.0E+00 O14867			0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00					0.0E+00	0.0E+00	0.0E+00 A	0.0E+00		0.0E+00	0.0E+00	0.0E+00 A
Expression Signal	1.91	1.09	1	0.83	0.83	1.01	1.01	4.46	4.46	1.21	6.0	1.09	4.19	7.89	75.43	1.39	1.39	86.0	2.82	2.36	2.36	1.15	1.16	1.16	1.16	1.18	1.17	1.26	36.36	2.74	1.32	1.1
ORF SEQ ID NO:	13616	13618	13620	13622		13632	13633	13635	13636	13657	13658	13662	13664		13685	13689	13690		13731	13735			13742			13747		13748	13750	13752	13753	13754
Exan SEQ ID NO:	8098	8810	8612	8614		8625				8651	8652	8656	6998	П	8682	2898	8687	8733	8734	8737			8742	8745		8746	8747	8748	8751	8753		8756
Probe SEQ ID NO:	3601	3603	3605	3607	3607	3618	3618	3623	3623	3645	3646	3650	3653	3674	3677	3683	3683	3729	3730	3733	3733	3734	3738	3741	3741	3742	3743	3744	3747	3749	3750	3752

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אוונופופ באחון ביוחקס באליו המספר וויין ביין ביין ביין ביין ביין ביין ביי	Top Hit Descriptor Database Source											T HUMAN	NT Hamo sapiens inocome protein 50 (1/1 05), milkers.  Hamo sapiens cDNA clone DKFZp434N0413 6 1 424 (supported the sapiens cDNA clone DKFZp434N0413 6 1	HOMAN								EST HUMAN 601236966F1 NIH MGC 44 Home sapiens CON Constitution of the constitution of	EST HUMAN 601193827F1 NIH MGC 7 Homo sapiens cDNA clone IMAGE:3537774 5	Т	T								
	₽ .	mo sapiens myosin light chain kinase isoform 2	mo sapiens 5-hydroxydyptamine (serotonin) rec	mo saplens transient receptor potential channel	mo saciens chromosome X open reading frame	mo sepiens chromosome X open reading frame	men zinc finger protein ZNF134 mRNA, compl	mo sapiens intersectin short isoform (ITSN) m		mo sapiens potassium voltage gated channel,	And sapielis coco il manage il muse sapiens d	43340 R28830 2. ;contains element PTR7 rep	umo sapiens (bosoma) protein 30 (n. 30), iii.	Services AP1 comme subunit binding prote	and services AP1 damma subunit binding prote	omo sapiens glutamate receptor, metabotropic	omo sapiens melanoma antigen, family B, 1 (M	omo sapiens HBP17 heparin-binding and FGF	omo sapiens ryanodine receptor 3 (RYR3) mR	omo sapiens zinc finger protein (KIAA0412) m	Iomo sapiens F-box protein Fbl3b (FBL3b) mix	01236966F1 NIH MGC 44 Homo sapiens CD	01193827F1 NIH MGC 7 Homo sapiens cDN	01193827F1 NIH_MGC_7 Homo sepiens cDN	Jomo saplens cancer-testis antigen CT10 (CT1	lomo sapiens canoer-testis entigen CT10 (CT1	Juman MHC class II lymphocyte entigen DPw4	Homo sapiens chromosome 21 segment HS21	Novel human gene mapping to chomosome 20	Homo saplens chromosome 21 segment HS21	Homo sapiens chromosome 21 segment HS21	Homo saplens eukaryotic translation elongetion	
XOII LIONS												T HUMAN		HOMAN									Т	Т	1 2 2								
Sirigie	Top Hit Acession No.	TN C POSCOS	0.0E+00 AF0099011.2	TIM 3050400	TIN 07 10021	4503176 NT	TN 4 24 24 24 24 24 24 24 24 24 24 24 24 2	-	T	4826783 NT	4759171 NT	1864727.1	9745	0.0E+00 AL040338.1 ES	6005887 N I	E005887 IN THE	TN 8505024	NE440442 4	36758		AF12953	BE378602.1	AW580740.1	BE264998.1	AE-204990.1	AF110193.1	AF110185.1	M23910.1	AL 103303.2	AL118494.1	AL163284.2	AL103200.4	
									• 1									_1_	_				-	919	818	318	힑	۶۱۶	< 1 ?	취	Ž١٩	ŽΙ	
	Aost Similar (Top) Hit BLAST E Value		0.0=+00	0.0E+00	0.0=+00	0.01+00	0.00=+00	0.05+00	0.00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.05+00	0.05-00	1	1	1_			╛								0.0E+00	
	<del></del>								1.26 0.05100	1.15 0.0E+00	2.04 0.0E+00	2.48 0.0E+00/		1.47 0.0E+00					0.87 U.UE+001		L		2.13 0.0E+0	1.01	1.01	4.2	4.2		5.99	0.93	2.94	2.2	113.41
	Expression Signal			1.03		7.3	7.3	4.10		13777	13780 2.04	13781 2.48	13795 20.27	13797	13803 0.97	13804 0.97	13805 1.95	1.87	13808 0.87	13817	13821 2.31	13838 3.61	13844 2.13	13845 1.01	13846 1.01	13872 4.2	13873 4.2	4.42	5.99	13887 0.93	13889 2.94	13897 2.2	113.41
	D Expression Signal		==	1.03	0.81	13770 7.3	13771 7.3	8769 13774 4.16	1.20	1.15	8776 13780 2.04	2.48	13795 20.27	1.47	13803 0.97	8797 13804 0.97	6798 13805 1.95	1.87	8803 13808 0.87	8811 13817 1.081	8815 13821 2.31	3.61	8838 13844 2.13	8839 13845 1.01	8839 13846 1.01	4.2	8869 13873 4.2		5.99	8888 13887 0.93	8891 13889 2.94	8899 13897 2.2	

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Table 4
Single Exon Probes Expressed in HBL100

													-					11-				4	-				<u> </u>	_~	- 1		- 14	-
Single Exon Probes Expressed in MbL1W	Top Hit Descriptor	tt55g08.xt NCI_CGAP_GC8 Homo septens cDNA clone IMAGE:2244734 3' striiter to TR:080309 060309 KIAA0563 PROTEIN.;	Human zho finger protein ZNF133	Chlorocebus aethiops mRNA for ribosomid protein S4X, complete ods	Homo sapiens mRNA for UGA suppressor (RNA-associated entigenic protein (tRNA48 gene)	Homo sapiens mRNA for KIAA0316 protein, partial cds	Homo sapiens chromosome 21 segment HS21C003	Homo sapiens mRNA for rape-2 (rape gene)	Homo sepiens mRNA for rape-2 (rapa gene)	Homo seplens retinoblastome-binding protein 4 (RBBP4) mKNA	Homo sepiens retinoblastoma-binding protein 4 (RBBP4) mRNA	Homo sapiens phosphoribosy/glychamide formyltransferase, phosphoribosy/glychamide synthefase, phosphoribosy/aminolmidazole synthefase (GART) mRNA	Homo sapiens G protein-coupled receptor 21 (GPR21), mRNA	Homo sapiens mRNA for KIAA0287 gene, partial cds	Homo sapiens ras GTPase activating protein-like (NGAP) mRNA	Homo sapiens IMP (inosine monophosphate) dehydrogenase 1 (IMPORT), mKNA	Novel human mRNA from chromosome 1, which has similarities to BAT2 genes	2s55e09.rf Soares retina N2b4HR Homo sepiens cDNA clone IMAGE:362920 5' similar to contains Alu renetitive element:	Homo saplens DGCR8 (DGCR8) mRNA, complete ods	Homo sapiens DNA polymerase zeta catalytic subunit (REV3) mRNA, complete cds	Homo sapiens protein kinase, X-linked (PRKO) mRNA			nft (60kD) (GABPA), mrthA	Homo sepiens hypothetical protein FLJ10379 (FLJ10379), mKNA	Homo sepiens hypothetical protein FLJ10379 (FLJ10379), mRNA	Homo sapiens mRNA for KIAA0895 protein, partiel cds		4GE:2010970 S		MRT4410707-100500-001-602 FLIO707 From September CONTA	
Exon Prope	Top Hit Database Source	EST_HUMAN	NT	NT	NT	NT	N	LZ	Į.	뒫	NT	<del>_</del>	LN	IN	NT	NT	NT	CCT HIMAN	TA	IZ	Ę	Z	N	NT	IN	ΤN	Ę	EST HUMAN	EST_HUMAN	EST_HUMAN	EST HUMAN	ESI HOMAN
Single	Top Hit Acesslon No.	Al657078.1	U09366.1	AB015610.1					AJ277278.1	5032026 NT	5032026 NT	4503914 NT	4885306 NT	AB006625.1	4758807 NT	11419297 NT	AL096857.1	A A040076 4	T		26947	4828947 NT	4503854 NT	4503854 NT	TN 1622391	R922391 NT	AB020702.1	AI982597.1	A1982597.1	BE184856.1	BE184856.1	BE274217.1
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00		0.0E+00				0.0E+00	0.0E+00	0.0E+00			0.0E+00	0.0E+00	0.0E+00/			00-100 00		0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	-			0.0E+00	0.0E+00	0.0E+00
	Expression Signal	1.34	2.58	12.81	3.58	1.39	1.76	2.58	2.58	14.88	14.88	1.91	4.78	1.41	9.0	7.12	3.68	3	0.90	200	444	111	1.04	1.04	1.39	1.39		6.13		2.24	2.24	10.6
	ORF SEQ ID NO:		13910	13925		13938	13939	13940	13941	13948	13949	13957			13961	13962			OF CO.					L							14001	
	Exan SEQ ID NO:	8914	L	8932	8939	8949	8950	L	8951	8929	8959	8968		1_	L	L		Ŀ		4000	_\				L			L	1	1		9014
	Probe SEQ ID NO:	3914	3916	3933	3941	3951	3952	3953	3953	3961	3961	3969	3973	3974	3977	3978	3979		3980	3880	3991	2006	4000	4000	4003	4003	4008	4012	4012	4014	4014	4018

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Table 4
Single Exon Probes Expressed in HBL100

		nine-gamma-glutamyltransferase) (TGM3)		9900095 3' straiter to SW:THIZ_BOVIN	14 April 1446 OE :3063117 FT	IA TOTAL SUBSTITUTION OF THE O	Ą	sperm receptor for egg jelly, sea urchin	4GE:743197 3' similar to contains Alu	AGE:743197 3' similar to contains Alu		IKNA, complete cds	0								3	sapiens cDNA clone (MAGE:1724579 3'	Hand I	orax (Drosophila) homolog); translocated to, 4		3 41		
Single Exon Propes Expressed in HBL100	Top Hit Descriptor	Homo sepiens transglutaminese 3 (E potypeptide, protein-glutamine-gamma-glutamytransferase) (TGM3) mRNA	Homo sapiens nuclear receptor coactivator 3 (NCOA3), mRNA	De51704.x1 NIH_MGC_10 Homo explens cDNA done INACE:2900095 3' similar to SW:THIZ_BOVIN Qe5108 MITOCHONDRIAL THIOREFLOXIN PREFCI IS CAD.	UI-HF-BMO-adx-C-02-0-UI-J NIH MGC 38 Homo services CDNA closes MACE 3063147 5	Homo saplens hypothetical protein FL/10498 (FL/10498) mRNA	Homo saplens hypothetical protein FLJ10498 (FLJ10498), mRNA	Homo saptens polycystic ledney disease (polycystin) and REJ (sperm receptor for egg jelly, sea urchin homolog)-like (PKDREJ) mRNA	2468h07.s1 Sogres_testis_NHT Homo sapiens cDNA clone INAGE:743197 3' similar to contains Au repetitive element.contains element MFR35 repetitive element.contains element MFR35 repetitive element.	zu68h07.s1 Soares_testis_NHT Homo sepiens GNA clone IMAGE:743197 3' similar to contains Alu	Homo seriore DNA notweets with the transfer of	Home control of the Parking and Calaburg Submit (NEVS) michael, complete cas	Home seriors title (TTN) mRNA	Homo sepiens KIAA0440 protein (KIAA0440) mRNA	Homo sapiens desmoplakin (DPI) (DSP) mRNA	Homo sapiens desmoplakin (DPI, DPII) (DSP) mRNA	Homo sapiens chromosome 21 segment HS21C103	Homo sepiens mRNA for offactory receptor protein, pseudogene	Human apolipoprotein B-100 mRNA, complete cds	PM2-DT0023-080300-004-a08 DT0023 Homo sepiens cDNA	Homo sapiens F-box protein Fb/4 (FBL4) mRNA, pertial cds	qd23f06.x1 Soeres_placenta_8b3weeks_2NbHP8to8W Homo sapiens cDNA clone IMAGE:1724579.3* similar to contains MER20.b2 MER20 repetitive element:	Human CBFA3 (Cbfa3) gene, partial cds	Homo sepiens myeloid/lymphoid or mixed-lineage leukemia (trithorax (Drocophila) homolog); translocated to, 4 (MLLT4) mRNA	Homo seplens protein kinase C, nu (PRKCN), mRNA	Homo sepiens protein kinase C, ru (PRKCN), mRNA	Human G2 protein mRNA, partial cds	Human G2 protein mRNA, partial cds
e Exori Prope	Top Hit Database Source	_ <u></u> \	N	EST HUMAN	EST HUMAN	Z	NT	_ <b>t</b>	EST HUMAN		Т									EST HUMAN		EST HUMAN	П				ΤN	TN
Billo	Top Hit Acessian No.	4507478 NT	5729725 NT	0.0E+00 AW675599.1	0.0E+00 AW408788.1	8922466 NT	8922466 NT	5174632 NT	0.0E+00 AA401438.1		T	07720	4507720 IN	7662125 NT	4758199 NT	4758199 NT	163303.2	1.1			0.0E+00 AF174590.1	0.0E+00 AI189844.1		6174574 NT	6563384 NT	6563384 NT	10991.1	10991.1
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.05	0.0E+00	00E+00	005-00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 AI	0.0E+00	0.0E+00 J02610.1	0.0E+00	0.0E+00/	0.0E+00/	0.0E+00 U14520.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 U
	Expression Signal	1.02	2.26	6.78	0.95	1.51	1.51	2.18	9.69	og a	1.32	0 0	0.94	1.45	1.28	1.28	0.71	1.99	2.21	0.86	4.94	3.4	5.07	0.82	1.19	1.19	1.28	1.28
	ORF SEQ ID NO:		14008		14021	14024	14025		14049	14050	14058	14068	14069	14072	14079	14080		14113	14130	14146	14153			14160	14173	14174	14180	14181
	Exan SEQ ID NO:		9021	8028			. 9035	9044	9062	8062			L	1808	9095	9095	9103	9130	9146	9459	9166	9171	9173	9176	9191	9191	9198	9198
	Probe SEQ ID NO:	4024	4025	4032	4037	4039	4039	4048	4068	4068	4072	4085	4085	4087	4101	4101	4109	4135	4151	4164	417	4177	4180	4183	4198	4198	4205	4205

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Table 4
Single Exon Probes Expressed in

		_	_		_	-		_	_	_	_	,	_	<del></del>		_	_	_	-	<u> </u>	8	<u> </u>	<b>U.P.</b>		-	<u>B.</u>	<del>/</del>	<u></u>	<b>8</b>		
Single Exoli Plobes Expressed in ABL100	Top Hit Descriptor	Homo saplens COMPLEMENT COMPONENT C1a RECEPTOR (C10R) mRNA	Homo septens gap junction protein connectin 36 (C)(36) gane, complete cds	Homo sapiens plasma membrane calcium ATP ase isoform 1 (ATP2B1) gene, elternative spiloe products, partial cds	H.sapiens H2Bfh gene	H.sepkens H2Bih gene	xg68e10.x1 NCI_CGAP_UM Homo sapiens CDNA clone IMAGE.2833514.3' similar to TR.P97365 P97365 ZINC FINGER PROTEIN 84:	H.saplens H4/d gene for H4 histone	H. saplens H4/d gene for H4 histone	Homo sapiens KIAA0390 gene product (KIAA0390), mRNA	Homo sapiens KIAA0390 gene product (KIAA0390), mRNA	Homo sapiens caudal type homeo box transcription factor 4 (CDX4), mRNA	Homo sapiens Xq pseudoautosomal region; segment 2/2	Homo sapiens chromosome 21 segment HS210007	Homo sapiens myosin regulatory light chain interacting protein (MIR), mRNA	Homo saplens membrane bound aminopeptidase P (XNPEP2) gene, complete cds	Homo sapiens ACTN2 gene for alpha-Actinin 2, exon 10	Homo sapiens ACTN2 gene for alpha-Actinin 2, exon 10	24g7 Human retina cDNA randomly primed sublibrary Homo sapiens cDNA	24g7 Human retina cDNA randomly primed sublibrary Homo sapiens cDNA	Homo sapiens HPS1 gene, Intron 5	Human endogenous retrovirus HERV-K10	xx58e08.X1 NCL_CGAP_Esc2 Homo septiens cDNA clone IMAGE:2589446 3' similiar to SW:AHNK_HUMAN 009566 NEUROBLAST DIFFERENTIATION ASSOCIATED PROTEIN AHNAK;	Homo sapiens LIM domain kinase 2 (LIMK2), transcript variant 2a, mRNA	Homo sapiens vascular endothelial cell growth factor 165 receptor/meuropilin (VEGF165) mRNA, complete cds	Homo sapiens chromosome 21 segment HS21C007	PM1+HT0305-101199-002-d03 HT0305 Homo sapiens cDNA	Homo sapiens mRNA for putative ankyrin-repeat containing protein (ORF1)	Homo saplens mRNA for putative ankyrin-repeat containing protein (ORF1)	Homo sapiens G protein-coupled receptor 50 (GPR50) mRNA	Homo saplens serine-threonine protein kinase (MNBH) mRNA, complete cds
TYOU LION	Top Hit Database Source	Į,	F	<b>\</b>	Z	Z	EST_HUMAN	N	F	7	5	5	FZ	NT	5	NT	TN	77	EST_HUMAN	EST_HUMAN	NT	NT	EST_HUMAN	누	<u></u> -₽	Ę	EST_HUMAN	NT	NT	T.	N.
	Top Hit Acession No.	6912281 NT	AF153047.2	L14561.1			AW166933.1		X60483.1	7662091 NT	7662091 NT	4885126 NT	AJ271736.1	AL163207.2	· 7019456 NT	AF195953.1	AJ249765.1	AJ249765.1			AF200629.1	M14123.1	AW084964.1	8051619 NT	AF016050.1	AL163207.2	AW381570.1	AJ278120.1	4.1278120.1	58467	AF108830.1
	Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	_	-	0.0E+00	_	_	0.0E+00 /	0.0E+00	0.0E+00/	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00/		_	0.0E+00
	Expression Signal	10.08	1,05	5.12	4.71	4.71	1.5	2.06	2.06	11.18	11.18	11.95	1.08	0.99	1.08	6.33	2.96	2.96	0.81	0.81	2.03	0.76	2.57	1.15	0.91	7.59	1.24	1.16	1.16	1.24	3.1
	ORF SEQ ID NO:	14187		14222	14227	14228	14229	14235	14236		14243	14258	14259		14286		14299	14300	14306	14307			14344	٠	14346		14350	14357	14358	14360	14361
	Exan SEQ ID NO:		9229	9239	9243	9243	9244	9250	9250	9254	9254	9267	9268	9269	9301	9312	9317	9317	9321	9321	9335	9353	9364	10052	8366	6926	9371	9377	9377	9379	9380
	Probe SEQ ID NO:	4215	4235	4246	4249	4249	4250	4256	4256	4261	4261	4274	4275	4276	4309	4320	4325	4325	4330	4330	4344	4362	4372	4374	4375	4378	4380	4386	4386	4388	4389

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Table 4
Single Exon Probes Expressed in HBL100

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Top Hit Descriptor		Homo sepiens ATP-sensitive inwardly rectifying K-chararel subunit (KCNJ6/BIR1) gene, exon	Hamo sapiens pyrin (MEFV) gene, camplete cds	Homo sepiens pyrin (MEFV) gane, complete cds	Homo sapiens zino finger protein 195 (ZNF195), mRNA	Homo seplens syncytin precursor, mRNA, complete cds	Homo sapiens protocadherin gamma C3 (PCDH-gamma-C3) mRNA, complete cds	Homo saplens zinc finger protein 211 (ZNF211), mRNA	Homo sapiens eukaryotic translation elongation factor 1 alpha 1 (EEF1A1) mRNA	Homo saplens low density (boprotein receptor-related protein 6 (LRPs) mRNA, and translated products	Homo sepiens chondroitin sulfate proteoglycan 4 (melanome-associated) (CSPG4) mRNA	Homo saplens calcium/calmodulin-dependent protein kinase IV (CAMK4) mRNA	601447932F1 NIH_MGC_65 Home saplens cDNA clone IMAGE:3852127 5	Homo sapiens iduronate sulphate sulphatase (IDS) gene, complete cds	Homo sapiens KIAA0390 gene product (KIAA0390), mRNA	Homo sapiens KIAA0390 gene product (KIAA0399), mRNA	Homo sapiens PTEN (PTEN) gene, exons 3 through 5	Homo sepiens mRNA for G7c protein (G7c gene located in the class III region of the major histocompetibility complex)	Homo saplens mRNA for G7c protein (G7c gene located in the class III region of the major histocompatibility complex)	Homo saciens mRNA for KIAA0795 protein, partial cds	2018g08.s1 Strategene fetal retina 937202 Homo sapiens cDNA clone IMAGE:6098543'	Homo sapiens odz (odd Ozken-m, Drosophila) homolog 1 (ODZ1), mRNA	Homo sapiens chromosome 21 segment HS21C084	Homo sapiens cyclophilin-related protein (NKTR) gene, complete cds	Homo sapiens chromosome 21 segment HS21C100	Homo sapiens gene for natriuretic protein, partial cds	Homo sapiens DNA mismatch repair protein (MLH3) gene, complete cds	Homo sapiens mRNA for KIAA0406 protein, partiel cds	Homo saplens keratin 18 (KRT18) mRNA	Homo saplens keratin 18 (KRT18) mRNA	Mus musculus E-cadherin binding protein E7 mRNA, complete cds	Homo sapiens ADP/ATP carrier protein (ANT-2) gene, complete cds
Top Hit Database	Source	TN	IN	NT	TN	Z	LN.	LX.	NT	_ <b>⊦</b> N	TN	NT	T_HUMAN	NT.	FX	NT	TN	- FV	<u>_</u>	N N	EST_HUMAN		NT	NŢ	NT	TN		F	TN			FZ.
Top Hit Acession No.		S78684.1	0.0E+00 AF111163.1	0.0E+00 AF111163.1	TN 8265009	0.0E+00 AF208161.1	0.0E+00 AF152337.1	5454175 NT	4503470 NT	4505016 NT	4503098 NT	4502556 NT	E871908.1	15485.1	7662091 NT	7662091 NT	4F143314.1	1,245418.1	0.0E+00 A.1245418 1		0.0E+00 AA174072.1	7657410 NT	.163284.2		0.0E+00 AL163300.2			0.0E+00 AB007866.2	4557887 NT	4557887 NT	-167441.1	78810.1
Most Similar (Top) Hit BLAST E	Value	0.0E+00 S78684.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 BI	0.0E+00 L3	0.0E+00	0.0E+00	0.0E+00 A	0.0E+00 A.	0.05+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 AL	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00/	0.0E+00	0.0E+00	0.0E+00 AI	0.0E+00 L.
Expression Signal		1.05	1.06	1.06	3.16	5.97	1.83	1.07	50.75	1.01	1.5	1.11	1.16	2.72	11.71	11.71	2.89	10.04	10.04	0.8	1.45	1.39	2.8	1.18	5.1	1.83	6.0	1.33	11.33	11.33	2.06	0.94
ORF SEQ ID NO:					14384			14399	14407	14412	14415	14420			14423	14424	14440	14442	14443	1444		-		14463	14464		14465	14469	14475	14476	14477	14489
Econ SEQ ID	•			J	10053			9411	9421	9428	9432	9436	9439	9442	9444	9444	9461	9463	9463	9465	9480	9482	9484	9485	9486	9487	9489	9492	9497	9497	9498	9208
Probe SEQ ID	<u>i</u>	4398	4389	4399	4409	4413	4418	4421	4431	4438	4442	4446	4449	4452	4454	4454	4471	4473	4473	4475	4490	4492	4494	4495	4496	4497	4499	4502	4507	4507	4508	4518

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Table 4

Single Exon Probes Expressed in HBL100

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Top Hit Descriptor	Homo septens ADP/ATP cerrier protein (ANT-2) cene, complete cds	Homo sepiens ADP/ATP certer protein (ANT-2) sene, complete cds	Homo septens mRNA for KIAA 1047 protein, partial cds	Homo saplens mRNA for KIAA1047 protein, partial ods	Human endogenous retrovirus type K (HERV-K), as and any sense	QV2-BT0635-160400-142-h05 BT0635 Homo seniens cTNA	zv96b07.s1 Soares NiHMPu_S1 Homo sepiens cDNA clone MAGE:7676053*	Homo sapiens truncated tenascin XB (TNXB) gene, partial cds and TNXA gene recombination breakpoint region	Homo sapiens mRNA for KIAA1399 protein, partial ods	Homo sapients mRNA for KIAA1399 protein, partial cyls	Human displacement protein (CCAAT) mRNA	Homo sapiens butyrophilin, subfamily 2, member A2 (BTN2A2), mRNA	Homo sapiens butyrophilin, subfamily 2, member A2 (BTN2A2) mRNA	601158935F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3505521 5	Homo sapiens ecotropic viral Integration site 28 (EVI2B), mRNA	Homo sapiens ecotropic viral integration site 2B (EVI2B), mRNA	Human AHNAK nucleoprotein mRNA, 5' end	Human haptoglobin and haptoglobin-related protein (HP and HPR) genes, complete cds	Human haptoglobin and haptoglobin-related protein (HP and HPR) genes, complete cds	Homo sapiens cyclophilin-related protein (NKTR) gene, complete cds	Homo sapiens KIAA1084 protein (KIAA1084), mRNA	Homo sapiens KIAA0563 gene product (KIAA0563), mRNA	Human proto-oncogene tyrosine-protein kinasa (ABL) gene, expn 1e and expns 2-10, complete cals	Novel human mRNA from chromosome 1, which has similarities to BAT2 genes	Human CYP2D7AP pseudogene for cytochrome P450 2D6	Homo sapiens alpha-3 type LX collagen (COLBA3) gene, promoter region, and exons 1-26	Homo sapiens G-protein coupled receptor (RE2), mRNA	Homo saplens G-protein coupled receptor (RE2), mRNA	Homo saplens proteinx0008 (AD013), mRNA	Homo saplens proteinx0008 (AD013), mRNA	UI-H-BI3-qiw-c-04-0-UI.s1 NCI_CGAP_Sub5 Homo sepiens cDNA clone IMAGE:2733294 9'	Homo sapians aldehyda dehydroganasa 12 (ALDH12) mRNA, compiate cds
Top Hit Database Source	FA.	ZZ.	Į.	N	LN L	EST HUMAN	EST_HUMAN	LN	Z E	₽ E	R	N <sub>T</sub>	N P	EST_HUMAN	TA	בּוֹ	NT	Ę	LZ	NT.	ΥT	トフ	_ <del>_</del> _	LN	H	NT	L7	<u>ار</u>	رحا حا		T_HUMAN	NT
Top Hit Acession No.	78810.1	78810.1	4B028970.1	4B028970.1	/18890.1	3E081527.1	0.0E+00 AA418246.1	0.0E+00 AF086641.1	Γ			453812	6453812 NT	3E278730.1	5729817 NT	5729817 NT				0.0E+00 AF184110.1	7662479 NT	7662181 NT		0.0E+00 AL096857.1	Γ	0.0E+00 AF026801.1	1N 0011199	6677700 NT	7019320 NT	3320	0.0E+00 AW444637.1	
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 Y18890.1	0.0E+00 B	0.0E+00	0.0E+00	0.0E+00/	0.0E+00/	0.0E+00 M74099.1	0.0E+00	0.0E+00	0.0E+00 B	0.0E+00	0.0E+00	0.0E+00 M	0.0E+00 M69197.1	0.0E+00 M69197.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00 U07563.1	0.0E+00	0.0E+00 X58467.1	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00 A
Expression Signal	9.0	0.84	1.58	1.58	3.45	1.32	1.01	2.02	2.53	2.53	2.25	2.86	2.86	1.17	1.12	1.12	5.79	2.21	2.21	1.86	0.68	2.97	1.07	1.24	1.15	1.22	1.69	1.69	0.75	0.75	1.65	1.01
ORF SEQ ID NO:				14483	14489	14507	14508		14517		14519		14522		14546	14547	14551	14554	14555	14558	14559	14560	14568	14572		14586	14588	14589	14590	14591	14618	14627
Exan SEQ ID NO:							9521	9256	9530	9530	9531			9537	9558	9558	9563	9266	9266	9569	9570	9571	9578	9582	9587	9600	9602	9602	9603	9603	9626	9632
Probe SEQ ID NO:	4518	4518	4519	4519	4524	4530	4531	4536	4541	4541	4542	4544	4544	4549	4570	4570	4575	4578	4578	4581	4582	4583	4590	4594	4599	4615	4617	4617	4618	4618	4641	4647

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Table 4
Single Exon Probes Expressed in HBL100

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Single Exon Plobes Expressed in 1951 co	Top Hit Descriptor	Homo septens HSPC024-tsp mRNA, complete ods	Human WHC class I transplentation antigen (f/e) gene	Human MHC class I transplantation entigen (hia) gene	1. Contract of the state of the state of (GSTT2) and guitathions S-transferase the state (GSTT1)	rigino seprens guarantere de cuantere de companya de companya de companya de complete de companya de c	M.fascicularis mRNA for metalloprotease-like, disintegrin-like protein, IVa	Homo sapiens Williams-Beuren syndrome deletion transcript 9 (WBSCX9) mKNA, complete cus	Mus musculus zinc finger transcription factor Kalso mRNA, complete ods	Homo sapiens fragile X mental retardation 2 (PMR2) mRNA	Homo sapiens actin, alpha, cardiac muscle (ACTC), mRNA	Homo saplens hypothetical protein DKFZp762E1312 (DKFZp762E1312), mkNA	Homo sapiens hypothetical protein FLJ20073 (FLJ20073), mRNA	Homo sapiens KIAA0187 gene product (KIAA0187), mRNA	Himman Tra-Cydelta gene. exons 1-4: Tor-V-delta gene, exons 1-2; T-cell receptor alpha (Tor-alpha) gene, J1-	Jet segments; and Tor-Calpha gene, exons 1-4	Human Tor-C-delta gene, exons 1-4; Tor-V-delta gene, exons 1-2; 1-cell receptor agins (1 or express gene, exons	J61 segments; and Tcr-C-alpha gene, exprs 1-4	H.saplens MeCF-2 gene	H. saplens MeCr-2 gare	Homo sapiens chromosome 21 segment noctions and feder RNA references II 1, 28(D (TAF2I)	Home sapiens I A I A box binding protein ( I br. ) resources (cont.)	H. saplens MICA gene	Homo saplens zinc finger protein (NAAU414) minum	Homo sapiens mRNA for KIAA1443 protein, perdal cos	Homo sepiens mRNA for KIAA0635 protein partial case	Mus musculus zinc finger protein interacting with N protein 1 (Zin I), III NYA	Homo sapiers meningioma expressed antigen 6 (colled-coll prome-rical) (wochol) in the	Homo septens desmoplekin (DPI, DPII) (DSP) mRNA	Homo saplens MHC class 1 region	Homo sepiens opioid receptor, delta 1 (OPRD1) mRNA	Homo sepiens splice vertent AKAP350 mRNA, pertial ods	Home septens COLAA6 gene for a6(IV) collagen, even 44 and partial cds	
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	Most Similar (Top) Hit BLAST E Value	A OF TOO	30.100	0.0E+00 J00191.1	0.0=+00 300181.1	7 00±400	0 0E+00 X87205.1	00H+00	0.05+00	00-100	00110	200	0.05	0.0E+00	0.0E+00	0 0F+00 M94081.1		0.0E+00	0.0E+00 X94628.1	0.0E+00	0.0E+00	0.05+00	0.0E+00				L	١	١					0.05+00
	Expression Signal	1 44	1	0.68	0.66	808	0000	4.77	£, ,	10.7	40.50	50.51	2.33	22	2.67	2	iĝ:	1.91	8.1	1.8	1.97	127	3.82	1 68					1.04			2.39	_	1.02
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	Probe SEQ ID NO:		4649	4688	4688	1	4694	4698	4700	4701	4702	4704	4706	4709	4712	į	4713	4743	1745	4745	2,5		4/2/4	4/32	4734	4735	473(	4737	4738	4740	4744	4746	4747	4760

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Table 4
Single Exon Probes Expressed in HBL100

		93' similar to	93' similar to	632 Q61632			3,								39140	39140	39140										
Top Hit Desoriptor	Homo sapiens famesyi diphosphate synthese (famesyi pyrophosphate synthetase, dimethylallytranstransferase, geranytranstransferase) (FDPS) mRNA	qh68d08.X1 Soures, fetal liver_spieen_1NFLS_S1 Homo sapiens cDNA done IMAGE:1849839 3' similar to SW:ATPN_BOVIN Q28852 ATP SYNTHASE G CHAIN, MITOCHONDRIAL.	qh68d08.x1 Soares_fetal_liver_spleen_1NFLS_S1.Homo septens cDNA clone IMAGE:1849839.3' similar to SW:ATPN_BOVIN Q28852.ATP SYNTHASE G CHAIN, MITOCHONDRIAL.	qm15i05.xf NG_CGAP_Lu5 Homo septens cDNA done INAGE:1881921 3' similar to TR:Q61632 Q61632 EN-21.ACZ FUSION PROTEIN ;	Homo saplens chromosome 21 segment HS21C084	Homo sapiens KIAA0806 gene product (KIAA0806), mRNA	zq66b06.s1 Stratagene neurospithalium (#937231) Homo sapiens cDNA clone IMAGE:846547 3	Human ribosomal protein L21 mRNA, complete cds	Human endogenous retroviral DNA (4-1), complete retroviral segment	601303729F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3638118 5'	Homo sapiens desmoplakin (DPI, DPII) (DSP) mRNA	Homo sapiens mRNA for KIAA1043 protein, partial cds	Homo sapiens hypothetical protein FLJ20477 (FLJ20477), mRNA	Homo sapiens hypothetical protein FLJ20477 (FLJ20477), mRNA	no14g09.s1 NCI_CGAP_Phe1 Homo sepiens cDNA clone IMAGE:11007043' shriiler to TR:E239140 E239140 SPALT PROTEIN;	no14g09.s1 NCI_CGAP_Phet Homo sepiens cDNA done IMAGE:1100704.3' sknikar to TR:E239140 E239140 SPALT PROTEIN:	no14g09.s1 NCI_CGAP_Phot Homo sapiens cDNA done IMAGE:1100704 3' similar to TR:E239140	E239140 SPALT PROTEIN;	Homo sapiens HSPC114 mRNA, complete ods	Homo sapiens HSPC114 mRNA, complete cds	Homo sapiens DNA mismatch repair protein (MLH3) gene, complete ods	Homo saplens E2F transcription factor 2 (E2F2) mRNA	Homo sapiens E6-AP ubiquitin-protein ligase (UBE3A) gene, exon 3	Homo sapiens MHC class 1 region	Homo sapiens chromosome 21 segment HS21C009	Homo sapiens gammma-cytoplasmic actin (ACTGP3) pseudogene	
Top Hit Database Source		EST HUMAN	Г		Г		LHUMAN	+ IN	П	EST_HUMAN 6		E E			EST HUMAN E	EST HUMAN	Т	EST_HUMAN E		-	F		F		TN	TX	
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Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00 A	0.0E+00	0.0E+00 A	0.0E+00,	0.0E+00	0.0E+00	0.0E+00 U14967.1	0.0E+00 M10976.1	0.0E+00	0.0E+00	0.0E+00/	0.0E+00	0.0E+00	0.0E+00	0.0E+00		0.0E+00	0.0E+00	0.05+00/	0.0E+00	0.0E+00	0.0E+00/	0.0E+00 U53588.1	0.0E+00	0.0E+00 D50657.1	
Expression Signal	1.96	1.13	1.13	1.07	1.2	1.02	0.86	6.67	1.6	2.99	3.55	2.16	2.53	2.53	1	-		<del>-</del>	1.59	1.59	0.81	0.92	1.28	1.27	1.11	27.33	170
ORF SEQ ID NO:	14732	14735	14736		14762	14787	14774		14789		14794	14797	14804	14805	14815	14816		14817	14818	14819	10289		14831	14833			2000,
Exan SEQ ID NO:	9746	9748	9748	9752	6776	9785	9792	9797	9807	6086	9813	9820	9830	9830	9843	9843		9843	9844	9844	5275	9849	9860	9862	9986	6986	2000
Probe SEQ ID NO:	4762	4764	4764	4768	4795	4801	4808	4813	4823	4825	4829	4836	4848	4848	4862	4862		4862	4864	4864	4867	4870	4881	4883	4887	4890	5,5,

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Table 4
Single Exon Probes Expressed in HBL100

Top Hit Descriptor	Hamo sepiens cuclorhilin (1984 CVP) mBNA	8992004.91 Strategies achies brain S44 Hours contant of the last of the same of the last of the same of the last of the same of the last of the same of the last of the same of the last of the same of the last of the same of the last of the same of the last of the same of the last of the same of the last of the same of the last of the same o	Homo society PR domain comfaming 4 with "Nic Januar, 1990 NAVA"	Human endocensus netrowins K   TR   15 and near nears	Homo sapiens SH2-containing professional Anguing Anguarda Anguarda	Homo sapiens K(AA0971 anders (KIAA0071)DNA	Homo saniens notessium voltane cated channel lot related formal.	Homo saniens serine-threoning periods (MNDL) was a serine-threoning period (MNDL) was a serine-threoning periods (MNDL) was a serine-threoning periods (MNDL) was a serine-threoning period (MNDL) was a serine-	Homo secient station through kinese (MNRH) minuth, complete cas	Homo sapiens acidic 82 kDa protein mRNA (HSI 115552) mRNA	Homo sapiens coequiation factor C (1 inulins polywhemis) homology/COCU/ mBNA	Homo saplens G-ordein coupled recentor (RE2) mRNA	Homo sapiens G-protein complet recentor (RE2) mRNA	Homo saciens mutt. (E. coli) homolog 3 (MI H3) mRNA	DKFZD434 2428 rf 434 (symmer Hess) Home series cDNA dose DKFZ-4341 2420 El
Top Hit Database Source		EST HUMAN			¥										EST HUMAN
Top Hit Acession No.	5454153 NT	AA683268.1	7362	Y08032.1	-	7662421 NT	4826795 NT	AF108830.1		7657203 NT	4758021 NT	TM 0077700	D877700 NT	7657336 NT	AL044081.1
Most Similar (Top) Hit BLAST E Value	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00	0.0E+00
Expression Signal	1.13	66.0	0.95	0.72	0.92	0.92	0.67	1.07	1.07	1.27	1.17	0.99	0.99	0.74	0.75
ORF SEQ ID NO:	14884	14891	14906	14912	14919	14933	14934	14940	14941	14955	14976	14989	14990	14993	15004
Exan SEQ ID NO:	2086	9913	8268	9934	9942	9955	9328	8963	9963	0866	10005	10020	10020	10024	10037
Probe SEQ ID NO:	4930	4936	4951	4957	4965	4980	4981	4990	4990	6009	5034	5049	5049	5053	2068

## CLAIMS

A spatially-addressable set of single exon nucleic acid probes for measuring gene expression in a sample derived
 from human Breast comprising a plurality single exon nucleic probes, said probes comprising any one of the nucleotide sequences set out in SEQ ID NOs: 1 - 5,074 or a complementary sequence, or a portion of such a sequence.

- 2. A spatially-addressable set of single exon nucleic acid probes as claimed in claim 1 wherein each of said plurality of probes is separately and addressably amplifiable.
- 3. A spatially-addressable set of single exon nucleic acid probes as claimed in claim 1 wherein each of said plurality of probes is separately and addressably isolatable from said plurality.
- 4. A spatially-addressable set of single exon nucleic acid 20 probes as claimed in any of claims 1 to 3 wherein said probes comprise any one of the nucleotide sequences set out in SEQ ID NOS.: 5,075 - 10,058.
- 5. A spatially-addressable set of single exon nucleic acid 25 probes as claimed in any of claims 1 to 4, wherein each of said plurality of probes is amplifiable using at least one common primer.
- 6. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 to 5 wherein the set comprises between 50 - 20,000 single exon nucleic acid probes.
- 7. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 to 6, wherein the

average length of the single exon nucleic acid probes is between 200 and 500 bp.

- 8. A spatially-addressable set of single exon nucleic acid 5 probes as claimed in any of claims 1 to 7, wherein at least 50% of said single exon nucleic acid probes lack prokaryotic and bacteriophage vector sequence.
- 9. A spatially-addressable set of single exon nucleic acid 10 probes as claimed in any of claims 1 to 8, wherein at least 50% of said single exon nucleic acid probes lack homopolymeric stretches of A or T.
- 10. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 9 characterised in that said set of probes is addressably disposed upon a substrate.
- 11. A spatially-addressable set of single exon nucleic acid
  20 probes as claimed in claim 10 wherein said substrate is
  selected from glass, amorphous silicon, crystalline silicon
  and plastic.
- 12. A microarray comprising a spatially addressable set of25 single exon nucleic acid probes as claimed in any of claims1 11.
- 13. A single exon nucleic acid probe for measuring human gene expression in a sample derived from human Breast

  30 comprising a nucleotide sequence as set out in any of SEQ ID NOs.: 1 5,074 or a complementary sequence or a fragment thereof wherein said probe hybridizes at high stringency to a nucleic acid molecule expressed in the human Breast.

14. A single exon nucleic acid probe as claimed in claim 13 comprising a nucleotide sequence as set out in any of SEQ ID NOs.: 5,075 - 10,058 or a complementary sequence or a fragment thereof.

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- 15. A single exon nucleic acid probe for measuring human gene expression in a sample derived from human Breast which is a nucleic acid molecule having a sequence encoding a peptide comprising a peptide sequence as set out in any of SEQ ID NOs.: 10,059 15,009, or a complementary sequence or a fragment thereof wherein said probe hybridizes at high stringency to a nucleic acid expressed in the human Breast.
- 16. A single exon nucleic acid probe as claimed in any one
  15 of claims 13 to 15 wherein said single exon nucleic acid
  probe comprises between 15 and 25 contiguous nucleotides of
  said SEQ ID NO.
- 17. A single exon nucleic acid probe as claimed in any one of claims 13 to 15, wherein said probe is between 3 25 kb in length.
  - 18. A single exon nucleic acid probe as claimed in any one of claims 13 17, wherein said probe is DNA, RNA or PNA.

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- 19. A single exon nucleic acid probe as claimed in any one of claims 13 18, wherein said probe is detectably labeled.
- 30 20. A single exon nucleic acid probe as claimed in any one of claims 13 19, wherein said probe lacks prokaryotic and bacteriophage vector sequence.
- 21. A single exon nucleic acid probe as claimed in any one 35 of claims 13 - 20, wherein said probe lacks homopolymeric

stretches of A or T.

22. A method of measuring gene expression in a sample derived from human Breast, comprising:

contacting the microarray of claim 12, with a first collection of detectably labeled nucleic acids, said first collection of nucleic acids derived from mRNA of human Breast; and then measuring the label detectably bound to each probe of said microarray.

23. A method of identifying exons in a eukaryotic genome, comprising:

algorithmically predicting at least one exon from genomic sequence of said eukaryote; and then dètecting specific hybridization of detectably labeled nucleic acids to a single exon probe,

wherein said detectably labeled nucleic acids are derived from mRNA from the Breast of said eukaryote, said probe is a single exon probe having a fragment identical in sequence to, or complementary in sequence to, said predicted exon, said probe is included within a microarray according to claim 12, and said fragment is selectively hybridizable at high stringency.

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24. A method of assigning exons to a single gene, comprising:

identifying a plurality of exons from genomic sequence according to the method of claim 23; and then

measuring the expression of each of said exons in a plurality of tissues and/or cell types using hybridization to single exon microarrays having a probe with said exon,

35 wherein a common pattern of expression of said exons in

said plurality of tissues and/or cell types indicates that the exons should be assigned to a single gene.

- 25. A nucleic acid sequence as set out in any of SEQ ID 5 NOs: 1 10,058 which encodes a peptide.
  - 26. A peptide encoded by a sequence as set out in any of SEQ ID Nos: 1 10,058.
- 10 27. A peptide comprising a sequence as set out in any of SEQ ID Nos: 10,059 15,009.

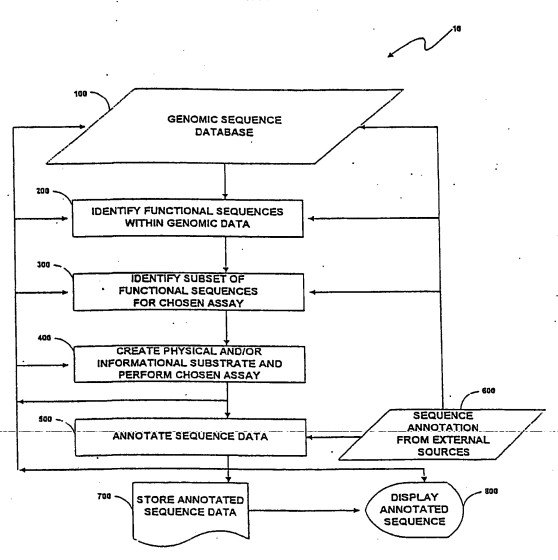


Fig. 1

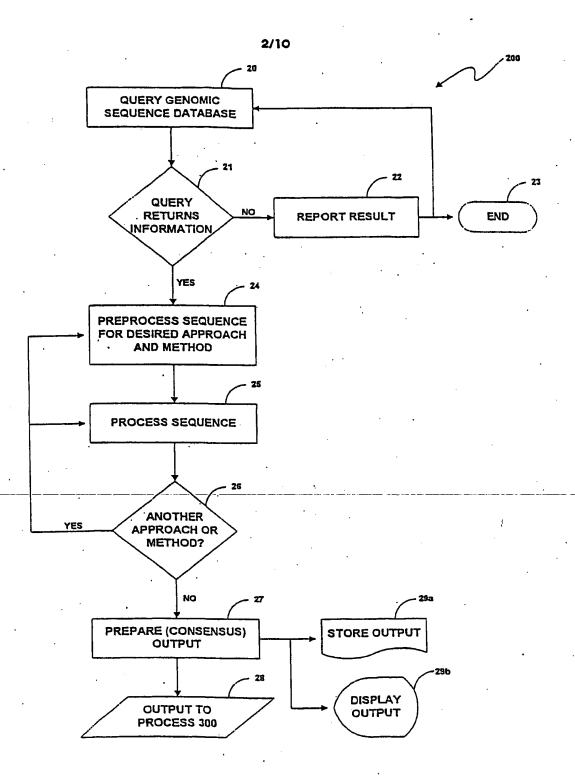


Fig. 2

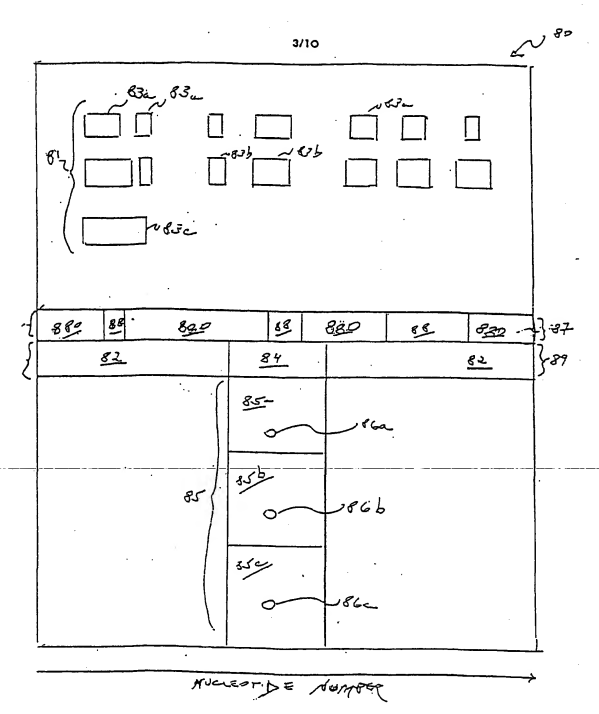


Fig. 3

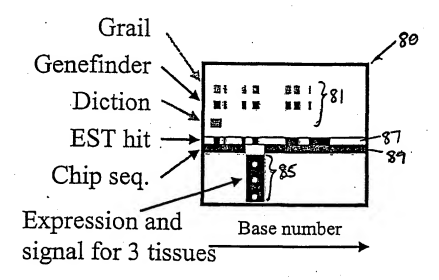


Fig. 4

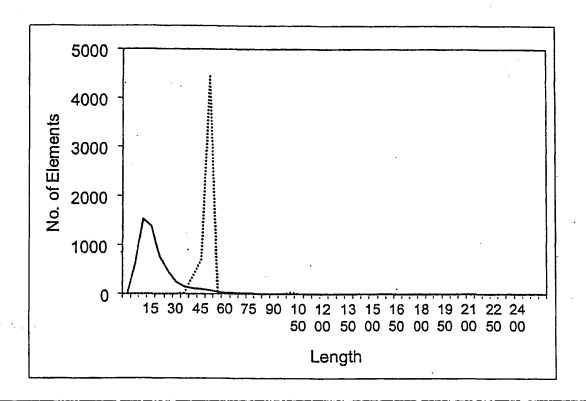


Fig. 5

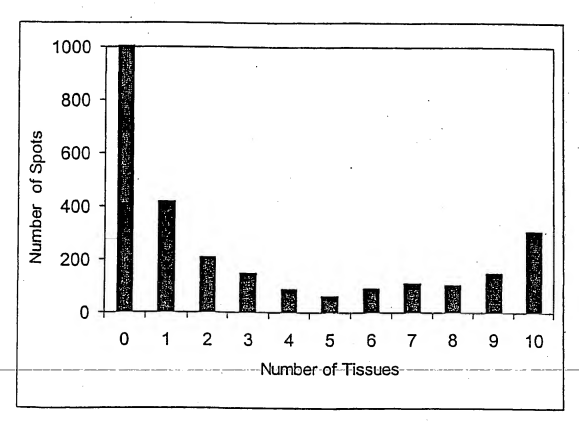
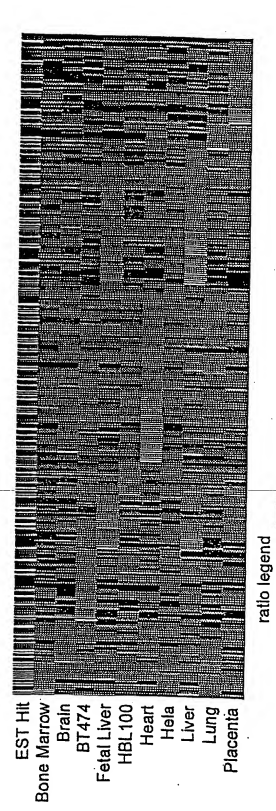


Fig. 6



ce-20 ce-60 ce-100

1 Fig. 7b

Fig. 7a

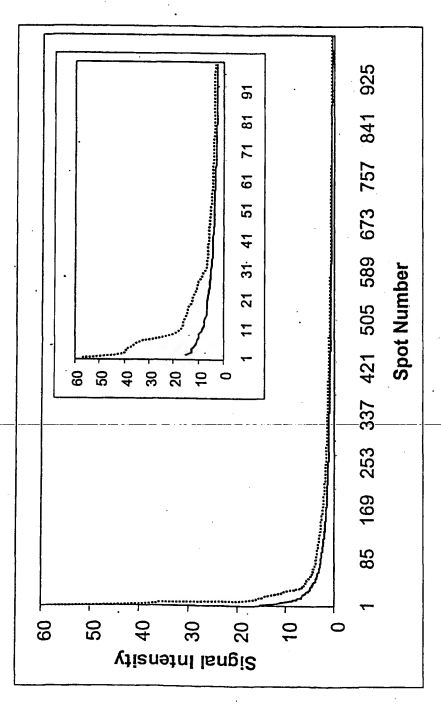


Fig. 8

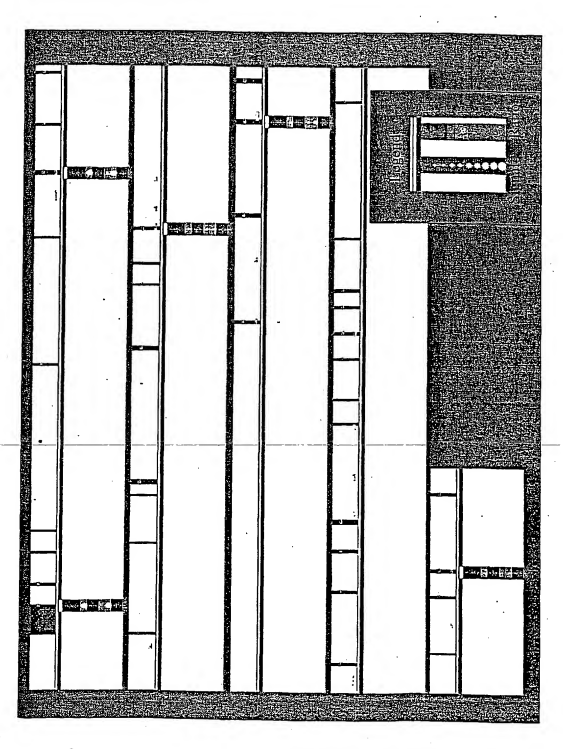


Fig. 9

Fig. 10

